AREA 317 RCRA QUARTERLY GROUND WATER MONITORING REPORT NO. 34 JANUARY THROUGH MARCH 1997

WHITTAKER CORPORATION, BERMITE FACILITY 22116 WEST SOLEDAD CANYON ROAD SANTA CLARITA, CALIFORNIA 91350 AME PROJECT NO. 21001.75

June 2, 1997

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21001.75/7

Subject:

Area 317 RCRA Quarterly Ground Water Monitoring Report No. 34

January through March 1997, Whittaker Corporation, Bermite Facility

22116 West Soledad Canyon Road, Santa Clarita, California

Dear Mr. Saebfar:

Enclosed is the Area 317 RCRA Quarterly Ground Water Monitoring Report No. 34 for the first quarter, January through March 1997. The monitoring was completed according to the requirements of the Water Quality Monitoring and Response Plan for the Interim Status Area 317 Surface Impoundment.

The statistical analyses for this sampling event showed that the established tolerance limits for pH, specific conductance, chloride, sulfate, TCE, TOC, or TOX were not exceeded. The tolerance limit for sodium was exceeded for the sample from monitoring well MW-10.

Please call me at (916) 939-7550 if there are any questions regarding the enclosed report.

Very truly yours,

ACTON • MICKELSON • ENVIRONMENTAL, INC.

Barbara J. Mickelson P.E.

California Registered Professional Engineer #43417

BJM:ecd Enclosure

cc/enc:

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AREA 317 RCRA QUARTERLY GROUND WATER MONITORING REPORT NO. 34 JANUARY THROUGH MARCH 1997

WHITTAKER CORPORATION, BERMITE FACILITY

TTAKER CORPORATION, BERMITE FACILITY 22116 WEST SOLEDAD CANYON ROAD SANTA CLARITA, CALIFORNIA 91350

1.0 INTRODUCTION

The Whittaker Corporation (Whittaker), Bermite facility (site) is located at 22116 West Soledad Canyon Road in Santa Clarita, California (Figure 1). Whittaker had interim status permits for 14 Resource Conservation and Recovery Act (RCRA) Hazardous Waste Management Units (HWMUs) when operations were terminated in April 1987. A document entitled "Whittaker Corporation, Bermite Division, Santa Clarita, California, CAD064573108, Facility Closure Plan Modifications" (Closure Plan), was prepared by Whittaker and approved by the California Environmental Protection Agency, Department of Toxic Substances Control (Cal-EPA) and U.S. Environmental Protection Agency (EPA) on December 28, 1987. The Closure Plan outlined procedures for obtaining approval by Cal-EPA and EPA of clean closure certification for the different HWMUs, including the 317 Surface Impoundment (Area 317).

The Closure Plan required the implementation of a ground water monitoring system at Area 317 capable of detecting and assessing the impact of the HWMU on the uppermost aquifer underlying Area 317. Implementation of a ground water monitoring system is described in the document entitled "Water Quality Monitoring and Response Plan for the Interim Status Area 317 Surface Impoundment," dated October 9, 1992 (Area 317 Monitoring Plan). This is a revised response plan approved by Cal-EPA which meets the requirements of the revisions to Title 22, California Code of Regulations and expands the constituents sampled and reported. The revised Area 317 Monitoring Plan was utilized for the nineteenth and subsequent sampling events.

A total of six ground water monitoring wells (MW-1, MW-3, MW-4, MW-5, MW-6, and MW-10) have been installed around Area 317 (Figure 2). Reports detailing the location and construction of monitoring wells, sampling and analysis plans for collecting and analyzing ground water samples from the ground water monitoring wells, abandonment of monitoring well MW-4, and quarterly sampling results which have been submitted to Cal-EPA and EPA are listed in Appendix A.

Quarterly ground water sampling activities were initiated on October 3, 1988, for monitoring wells MW-1, MW-3, and MW-4. The ground water monitoring program included analyses of water samples for volatile organic compounds (VOCs). Laboratory analytical results from the third quarterly sampling event reported trichloroethene (TCE) at 4,800 micrograms per liter (μ g/l) in the ground water sample collected from monitoring well MW-4. As a result, two additional monitoring wells were installed in Area 317 (MW-5 and MW-6).

The fourth quarterly monitoring event included sampling of the ground water from monitoring wells MW-1, MW-3, and MW-4. Monitoring wells MW-5 and MW-6 were not equipped for sampling during the fourth quarterly sampling event. Analytical results from the fourth quarter were similar to those reported in the third quarterly sampling event. The concentrations of VOCs reported in samples collected from monitoring wells MW-1 and MW-3 were below laboratory reporting limits; however, analysis of the ground water sample collected from monitoring well MW-4 reported TCE at 7,200 μ g/1. Analysis of ground water samples collected from monitoring well MW-4 during the fifth through twelfth quarterly sampling events reported a steady decline in TCE concentration. Based on the results of the initial four sampling events, a reduced list of chemical parameters, approved by Cal-EPA, was utilized for the fifth through eighteenth quarterly sampling events.

Statistical analysis of indicator parameters was initiated during the fifth quarterly sampling event. The ground water samples collected and analyzed for indicator parameters from monitoring wells MW-1, MW-3, and MW-4 for the initial year of monitoring were evaluated to assess whether statistically significant changes to the ground water had occurred as a result of site activities.

A Comprehensive Ground Water Monitoring Evaluation (CME) was conducted by Cal-EPA on January 24 and 25, 1990, during the sixth quarterly monitoring event. Personnel from Cal-EPA were present during all phases of the sixth quarterly monitoring event, from the taking of initial potentiometric surface elevation measurements to the sealing of the coolers containing the quarterly ground water samples.

For the thirty-fourth quarterly monitoring event, ground water samples from monitoring wells MW-1, MW-3, MW-5, MW-6, and MW-10 were collected on March 19, 1997. This report presents the results of the thirty-fourth quarterly ground water sampling event, along with recommendations for future quarterly ground water sampling events.

2.0 GROUND WATER LEVEL MEASUREMENTS

Water level measurements were taken on March 17, 1997, prior to well evacuation and sampling activities. Monitoring well locations with respect to Area 317 are shown on Figure 2. Water levels were measured to the nearest 0.01 foot. Water level elevations increased 0.74, 0.73, 0.84, 0.90, and 0.74 foot in monitoring wells MW-1, MW-3, MW-5, MW-6, and MW-10, respectively, between the thirty-third and thirty-fourth quarters. Table 1 summarizes potentiometric surface elevation data for monitoring wells in Area 317. Figure 3 illustrates the historic changes in potentiometric surface elevations in monitoring wells MW-1, MW-3, MW-5, MW-6, and MW-10. The March 17, 1997, water level measurements were used to develop the potentiometric surface contours illustrated on Figure 2. Figure 2 indicates that the inferred flow direction for March 17,

1997, is toward the north. Based on these data, monitoring wells MW-6 and MW-10 are estimated to be located hydraulically downgradient from Area 317, monitoring well MW-5 is estimated to be located hydraulically downgradient and/or crossgradient from Area 317, monitoring well MW-1 is estimated to be located hydraulically crossgradient and/or upgradient from Area 317, and monitoring well MW-3 is estimated to be located hydraulically upgradient from Area 317. The ground water flow direction estimated for March 17, 1997, is similar to the flow direction estimated for the previous sampling event. The estimated ground water flow direction has varied from north-northwest to north-northeast since initiating quarterly ground water monitoring, possibly contributing to the reported variability in ground water chemistry.

3.0 SAMPLE COLLECTION AND ANALYSES

Ground water evacuation, stabilization, and sampling procedures are tabulated in Appendix B. Included in these procedures are revisions requested by Cal-EPA representatives during a June 26, 1996, field visit to observe Bermite facility personnel conduct quarterly ground water sampling.

3.1 Required Ground Water Analyses

For the thirty-fourth sampling event, the following analytical parameters were tested according to the Area 317 Monitoring Plan:

 Ground Water Monitoring Parameters: pH, specific conductance, total organic carbon (TOC), total organic halogens (TOX), TCE, sulfate, sodium, manganese, iron, and chloride.

Background water quality parameters were not analyzed for the thirty-fourth sampling event based on the results from previous sampling events.

All ground water samples collected during the thirty-fourth sampling event were submitted to FGL Environmental (FGL) in Santa Paula, California. FGL Environmental is certified by Cal-EPA to perform the ground water analyses outlined in the Area 317 Monitoring Plan. Chain-of-custody and sample analyses request forms are included in Appendices C and D, respectively. A description of FGL's Quality Assurance/Quality Control (QA/QC) program is provided in Appendix E. Copies of the laboratory analytical reports for all trip, field, and method blanks, and duplicate and spiked samples analyzed by FGL are provided in Appendix F.

3.2 Approved Analytical Methods

Ground water monitoring parameters were analyzed by EPA or other approved methodologies.

Analytical methodologies were presented in the "Ground Water Sampling and Analysis Plan," dated August 1988. Modifications to this plan were approved by Cal-EPA prior to the fifth quarterly sampling event. Copies of the laboratory test method protocol were included in Appendix B of "Quarterly Sampling Report No. 1," dated December 1988.

A summary of sample volumes, sample containers, and laboratory analytical methods utilized during the thirty-fourth sampling event is presented in Table B-3, Appendix B. Procedures regarding sample containers, sample labeling, sample collection, and field QA/QC are tabulated in Appendix B.

4.0 GROUND WATER SAMPLE ANALYTICAL RESULTS

4.1 Ground Water Monitoring Parameters

Ground water samples from each monitoring well were analyzed for pH, specific conductance, chloride, iron, manganese, sodium, sulfate, TCE, TOC, and TOX to serve as ground water monitoring parameters. Table 2 summarizes the results of the ground water monitoring parameter analyses for the thirty-fourth sampling event, along with results from the previous four sampling events. Copies of the original laboratory reports are presented in Appendix G.

Laboratory pH measurements of 7.8, 7.6, 7.8, 7.7, and 7.8 were recorded for samples collected from monitoring wells MW-1, MW-3, MW-5, MW-6, and MW-10, respectively, for the thirty-fourth monitoring event. The laboratory pH measurements recorded for samples collected from the monitoring wells during the thirty-fourth sampling event are generally consistent with the measurements recorded during previous sampling events.

Specific conductance measurements of 770, 600, 530, 550, and 600 micromhos per centimeter squared (μ mhos/cm²) were recorded for samples collected from monitoring wells MW-1, MW-3, MW-5, MW-6, and MW-10, respectively, for the thirty-fourth sampling event. The specific conductance measurements recorded during the thirty-fourth sampling event are generally consistent with measurements recorded during previous sampling events. The results for chloride, sodium, and sulfate were 170, 53, and 10 milligrams per liter (mg/l) for the sample from monitoring well MW-1; 29, 60, and 71 mg/l for the sample from monitoring well MW-3; 54, 56, and 30 mg/l for the sample from monitoring well MW-5; 73, 56, and 29 mg/l for the sample from monitoring well MW-6; and 68, 86, and 38 mg/l for the sample from monitoring well MW-10. Detectable iron concentrations ranged from 60 μ g/l (MW-5) to 170 μ g/l (MW-6). The concentration of iron in the sample collected from well MW-3 was less than the 50 μ g/l detection limit. The results for chloride, sodium, sulfate, and iron are generally consistent with results from previous sampling events.

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The concentration of manganese in samples collected during the thirty-fourth sampling event ranged from less than the 0.5 μ g/l detection limit in the sample collected from monitoring well MW-3 to 3.2 μ g/l in the sample collected from monitoring well MW-1.

Laboratory analytical results for samples collected from Area 317 monitoring wells during the thirty-fourth sampling event did not indicate the presence of TCE, TOC, or TOX at method detection limits of 0.5 μ g/l, 0.5 mg/l, and 5 μ g/l, respectively. Trip blanks and field blanks prepared during sampling were also non-detect for TCE, TOC, and TOX. These analytical results are consistent with results from previous sampling events.

4.2 Background Water Quality Parameters

Background water quality parameters were not tested during the thirty-fourth monitoring event. The background water quality parameters were last tested during the twenty-third monitoring event because of a third consecutive exceedance with respect to the tolerance limit established for sodium. A summary of historical background water quality parameters is presented in Table 3.

5.0 STATISTICAL ANALYSIS OF RESULTS TO DATE

As was indicated in the document entitled "Ground Water Sampling and Analysis Plan," dated August 1988, and required in 40 CFR Part 265.92, statistical analysis of the indicator parameters was previously performed to determine whether a statistically significant difference in the water quality existed between the individual downgradient monitoring wells and the upgradient or background monitoring wells. At that time, monitoring wells MW-1 and MW-3 were considered upgradient or crossgradient relative to Area 317, and monitoring wells MW-5, MW-6, and MW-10 were considered downgradient or crossgradient relative to Area 317.

After four quarters of sampling and analysis of the monitoring system, the mean, standard deviation, variance, and coefficient of variance of the four indicator parameters were calculated. These values were reported to Cal-EPA in correspondence to Mr. Alan Sorsher, P.E., Cal-EPA, from Wenck Associates, Inc. (Wenck), dated October 25, 1989. The statistical analysis, presented in the fifth through tenth quarterly sampling reports, indicated only one statistically significant difference in water quality as determined by the indicator parameters. This was interpreted by Wenck to be caused by erroneous TOC results from the sixth quarter.

Since the approval of the Area 317 Monitoring Plan by Cal-EPA, the statistical comparison of analytical results for each downgradient monitoring well is made against the tolerance limits calculated from upgradient monitoring well results for the ten ground water monitoring parameters (chloride, sulfate, iron, manganese, sodium, TCE, TOC, TOX, specific conductance, and pH). The tolerance limits for the ground water monitoring parameters are updated at a minimum annually to include the latest analytical data.

Concentrations of the ground water monitoring parameters in the ground water samples collected from Area 317 monitoring wells for the thirty-fourth quarter are included in Table H-1, presented in Appendix H. A summary of the quarterly statistics for each background monitoring well and the tolerance limit calculations for the ground water monitoring parameters are presented in Appendix H, Tables H-2, H-3, and H-4. Graphical presentation of the statistical information is also included in Appendix H.

5.1 Assumptions Used in the Statistical Analysis

As recommended in the document entitled "RCRA Ground Water Monitoring Technical Enforcement Guidance Document," the data points that are less than the detection limit have been given a value equal to one-half the detection limit of the analyte. As recommended in the document entitled "Statistical Analysis of Ground Water Monitoring Data at RCRA Facilities, Interim Final Guidance" (Guidance Document), the statistical analysis assumes a value for the confidence coefficient (1-a) of 0.95 and a value for the proportion (y) of 0.95. This translates to a 95 percent confidence that 95 percent of future background monitoring well results will fall within the tolerance interval predicted. The tolerance limits for pH were calculated using a two-tailed distribution, and the tolerance limits for the other parameters were calculated using a one-tailed distribution. It was assumed that the data are distributed normally.

5.2 Data Preparation

The ground water sample analytical results from the two background or upgradient monitoring wells (MW-1 and MW-3) for all ground water sampling to date, and the results for the three downgradient monitoring wells (MW-5, MW-6, and MW-10) for the thirty-fourth quarter of ground water sampling, have been tabulated and prepared for statistical analysis. In accordance with the tolerance limit methodology used for this statistical analysis, the analytical results for the ten ground water monitoring parameters are summarized by quarter and by monitoring well.

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Arithmetic mean and standard deviation summary statistics have been calculated from background monitoring well results and are utilized in calculating the tolerance limits for each of the ground water monitoring parameters.

The statistical analysis for the ground water monitoring parameters requires a statistical comparison using tolerance intervals between water quality downgradient of Area 317 (wells MW-5, MW-6, and MW-10) and background water quality (wells MW-1 and MW-3). Tolerance intervals for each of the ten water quality parameters are based on the average of all the quarterly monitoring data for background wells MW-1 and MW-3. The most recent quarterly monitoring data for each downgradient well is compared to the established tolerance interval for each parameter to determine if an exceedance has occurred (Table H-1).

The calculations of the quarterly statistics were performed as outlined in the Area 317 Monitoring Plan. The values of K were taken from the statistical tables based on the number of samples and a one-sided tolerance limit. Note that pH values have not been reported as hydrogen ion concentrations as was done previously and that the value of K for the analysis of pH is derived from the tables for two-sided tolerance limits. TCE has never been reported above the detection limit in samples from monitoring wells MW-1 and MW-3; therefore, the tolerance limit for TCE is set at the detection limit.

5.3 Results

The thirty-fourth quarter results for each ground water monitoring parameter at each downgradient monitoring well were compared to the tolerance limits based on the first through thirty-fourth quarter results for background monitoring wells MW-1 and MW-3. The statistical analysis, summarized in Table H-1, indicates that there is no excursion of tolerance limits for pH, specific conductance, chloride, sulfate, iron, manganese, TCE, TOC, or TOX in downgradient ground water quality. The reported concentrations of sodium for monitoring wells MW-5 and MW-6 were below tolerance limits, but the analytical results for monitoring well MW-10 indicate a tolerance limit exceedance for sodium. In the past, an elevated sodium concentration in monitoring well MW-10 relative to sodium concentrations in the other four Area 317 wells has not indicated a statistical impact to ground water quality, based on the concentrations of the other ground water monitoring parameters and retesting of background water quality parameters.

6.0 SUMMARY OF RESULTS, JANUARY THROUGH MARCH 1997

6.1 Ground Water Level Measurements

The estimated direction of ground water flow based on the March 17, 1997 data is toward the north, which is consistent with the ground water flow direction estimated during the previous sampling event. Utilizing this data, monitoring wells MW-6 and MW-10 are estimated to be located hydraulically downgradient from Area 317; monitoring well MW-5 is estimated to be located hydraulically downgradient and/or crossgradient from Area 317; monitoring well MW-1 is estimated to be located hydraulically crossgradient and/or upgradient from Area 317; and monitoring well MW-3 is estimated to be located hydraulically upgradient from Area 317.

6.2 Ground Water Monitoring Parameters

The reported pH results for the thirty-fourth sampling event range from 7.6 for the sample from monitoring well MW-3 to 7.8 for samples from monitoring wells MW-1, MW-5, and MW-10. The specific conductance in samples from the five monitoring wells ranged from 530 μ mhos/cm² for monitoring well MW-5 to 770 μ mhos/cm² for monitoring well MW-1. Neither TOC nor TOX were detected in any of the samples collected at method detection limits of 0.5 mg/l and 5 μ g/l, respectively. The pH, specific conductance, TOC, and TOX results reported for the thirty-fourth sampling event are generally consistent with the results reported for the previous sampling events. The ground water sample analytical results for chloride, iron, manganese, sodium, and sulfates from the five monitoring wells are also generally consistent with historic data. The primary constituent of concern for Area 317, TCE, was not present in any of the samples collected during the thirty-fourth sampling event.

6.3 Background Water Quality Parameters

Background water quality parameters were not tested in this quarter. The Area 317 Monitoring Plan indicates that if tolerance limits for any water quality parameters are exceeded for three consecutive monitoring events, then additional samples will be collected during the subsequent monitoring event and analyzed for the background water quality parameters.

The tolerance limit for sodium was exceeded for the eleventh consecutive quarter in the sample from monitoring well MW-10. However, AME proposes that no further background water quality parameters be tested due to this exceedance at this time. Previous reports of elevated sodium concentrations in the samples from monitoring well MW-10 in comparison to concentrations for background wells has not indicated an impact with respect to the other ground water quality parameter concentrations. During the twenty-third sampling event, all six background parameters were analyzed due to the exceedance of the tolerance limit for sodium in the sample from

monitoring well MW-10. Analysis of the data indicated that the concentrations of background water quality parameters were consistent with the historical data for these parameters in samples from upgradient wells MW-1 and MW-3.

6.4 Statistical Analysis

The analytical results from the thirty-fourth quarter sampling event indicate that values for pH, specific conductance, chloride, sulfate, iron, manganese, TCE, TOC, and TOX in the downgradient monitoring wells are within the tolerance limits set by calculations using historical results from the background monitoring wells. The tolerance limit for sodium was exceeded for the sample from monitoring well MW-10, but this statistical exceedance does not appear to reflect an actual degradation of ground water quality.

7.0 RECOMMENDATIONS

The following recommendations are presented:

- Conduct future sampling events in accordance with the procedures set forth in the document entitled "Water Quality Monitoring and Response Plan for the Interim Status Area 317 Surface Impoundment," dated October 9, 1992.
- Update the tolerance limits for the ground water monitoring parameters following the thirty-fifth quarterly sampling event.

8.0 REMARKS

The recommendations contained in this report represent our professional opinions. These opinions are based on currently available information and were developed in accordance with currently accepted hydrogeologic and engineering practices at this time and location. Other than this, no warranty is implied or intended.

TABLE 1

POTENTIOMETRIC SURFACE ELEVATIONS RCRA GROUND WATER MONITORING WELLS WHITTAKER CORPORATION, BERMITE FACILITY

Well No.	MW-I	MW-3	MW-4	MW-5	MW-6	MW-10
Top of Casing Elevation	1,561.32	1,538.51	1,538.43	1,493.37	1,521.09	1,537,49
Date			Potentiometric	Surface Elevations		
12/23/87 01/27/88 02/03/88 02/03/88 02/04/88 02/05/88 02/09/88 02/10/88 02/11/88 02/12/88 02/12/88 04/05/88 04/05/88 04/05/88 04/12/88 04/12/88 04/12/88 01/23/89 04/17/89 07/27/89 08/10/89 08/18/89 01/24/90 04/16/90 07/16/90 07/16/90 01/28/91 04/22/91 07/17/90 10/28/91 10/08/91 10/08/91 10/08/91 10/08/91 10/08/92 01/25/93 06/07/93 09/20/93 12/06/93 03/07/94 06/21/94 09/13/94 12/12/94 03/27/95 06/26/95 09/08/95 12/04/95 03/18/96 06/24/96	1,107.81 1,108.03 1,108.36 1,108.36 1,108.36 1,108.24 1,108.28 1,108.28 1,108.11 1,107.69 1,107.76 1,107.66 1,107.56 1,107.61 1,107.86 1,107.56 1,107.86 1,109.82 1,097.37 1,094.67 1,093.93 1,093.62 1,092.07 1,090.56 1,088.66 1,088.56 1,071.22 1,063.63 1,055.22 1,071.22 1,063.63 1,055.22 1,071.22 1,063.63 1,055.22 1,071.22 1,063.63 1,055.22 1,051.88 1,050.47 1,046.84 1,043.87 1,044.79 1,049.24 1,052.40 1,057.58 1,056.22 1,057.58 1,056.22 1,057.58 1,056.22 1,053.94 1,054.62 1,057.58 1,056.22 1,053.94 1,054.62 1,057.58 1,056.22 1,053.94 1,064.21 1,068.60 1,066.792	1,109.51 1,109.88 1,109.14 1,109.17 1,109.13 1,109.27 1,108.86 1,108.23 1,107.62 1,097.62 1,094.85 1,094.89 1,093.76 1,092.16 1,090.54 1,083.78 1,070.54 1,071.29 1,063.78 1,076.54 1,071.29 1,063.78 1,055.41 1,055.29 1,050.88 1,047.40 1,044.58 1,045.61 1,050.36 1,054.11 1,056.27 1,059.63 1,058.38 1,056.25 1,056.79 1,061.45 1,062.97 1,063.59 1,066.41 1,070.81	1,102.61 1,100.77 1,098.92 1,096.05 1,093.53 1,092.64 1,091.08 1,087.83 1,082.29 1,078.86 1,075.46 1,069.75 1,061.66 1,053.28 1,050.63 1,049.33	1,093.02 1,092.32 1,092.03 1,090.62 1,089.17 1,087.23 1,081.41 1,078.25 1,074.64 1,068.90 1,060.53 1,052.12 1,049.76 1,048.78 1,045.14 1,042.05 1,044.22 1,049.19 1,052.47 1,052.47 1,052.47 1,055.69 1,055.41 1,052.79 1,055.61 1,060.35 1,060.35 1,061.06 1,064.15 1,068.75 1,065.99 1,065.71	1,093.15 1,092.49 1,092.19 1,090.64 1,089.50 1,087.32 1,081.85 1,078.56 1,074.91 1,069.25 1,061.14 1,052.69 1,050.06 1,048.92 1,042.13 1,043.64 1,048.70 1,051.79 1,053.58 1,054.93 1,052.44 1,053.55 1,059.28 1,059.28 1,060.66 1,063.65 1,067.99 1,065.54 1,067.28	1,050.57 1,049.37 1,045.77 1,042.77 1,044.29 1,052.53 1,054.53 1,057.77 1,055.86 1,053.43 1,054.50 1,059.89 1,066.82 1,068.99 1,066.61 1,068.99

^aNGVD = National Geodetic Vertical Datum. ^bMeasurement not recorded. ^cMonitoring well abandoned 05/28/92.

TABLE 2 GROUND WATER MONITORING PARAMETER ANALYTICAL RESULTS (MARCH 20, 1996 THROUGH MARCH 19, 1997)

Monitoring Well	Date	Specific Conductance (µmhos/cm²)	pH	TOC ^b (mg/l)	ΤΟΧ: (μg/l)	Chloride (mg/l)	Sulfate (mg/l)	TCE* (µg/l)	Iron (μg/l)	Manganese (μg/l)	Sodium (mg/l)
MW-1	03/20/96	770	7.4	< 0.5	<5	180	13	< 0.5	< 50	2.1	51
	06/26/96	760	7.7	< 0.5	<5	150	13	< 0.5	< 50	2.1	53
	09/19/96	750	7.6	< 0.5	7	160	12	<0.5	80	1.8	52
	12/05/96	750	7.5	< 0.5	<5	160	10	< 0.5	< 50	2.6	55
	03/19/97	770	7.8	<0.5	<5	170	10	<0.5	100	3.2	53
MW-3	03/20/96	610	7.6	< 0.5	<5	35	91	< 0.5	< 50	< 0.5	57
	06/26/96	600	7.9	< 0.5	<5	31	78	< 0.5	< 50	3.3	61
	09/19/96	610	8.0	< 0.5	<5	30	71	< 0.5	< 50	<0.5	56
	12/05/96	600	7.6	< 0.5	<5	26	62	< 0.5	< 50	<0.5	56
	03/19/97	600	7.6	< 0.5	<5	29	71	< 0.5	< 50	<0.5	60
MW-5	03/20/96	540	7.7	< 0.5	<5	57	36	< 0.5	< 50	1.6	55
	06/26/96	520	7.6	< 0.5	<5	50	34	< 0.5	< 50	1.5	55
	09/19/96	540	7.6	< 0.5	53	45	29	< 0.5	70	1.9	55
	12/05/96	520	7.7	<0.5	<5	52	30	< 0.5	< 50	1.6	60
	03/19/97	530	7.8	< 0.5	<5	54	30	<0.5	60	<2.5	56
MW-6	03/20/96	560	7.7	< 0.5	<5	76	29	< 0.5	< 50	1.4	53
1,1,1,	06/26/96	570	7.5	< 0.5	<5	70	34	< 0.5	<50	1.3	56
	10/01/96	570	7.6	< 0.5	8	76	28	< 0.5	100	1.2	48
	12/05/96	570	7.7	<0.5	<5	71	33	< 0.5	< 50	1.0	57
	03/19/97	550	7.7	<0.5	<5	73	29	<0.5	170	1.6	56
MW-10	03/20/96	610	7.8	< 0.5	<5	83	45	< 0.5	<50	1.7	82
1.2	06/28/96	600	7.7	< 0.5	<5	70	43	< 0.5	< 50	1.8	80
	09/25/96	610	7.6	< 0.5	<5	66	37	< 0.5	80	1.8	77
	12/05/96	610	8.1	< 0.5	<5	74	41	< 0.5	< 50	1.2	92
	03/19/97	600	7.8	< 0.5	<5	68	38	< 0.5	70	1.1	86

*TCE = Trichloroethene.

*TOC = Total organic carbon.

*TOX = Total organic halogens.

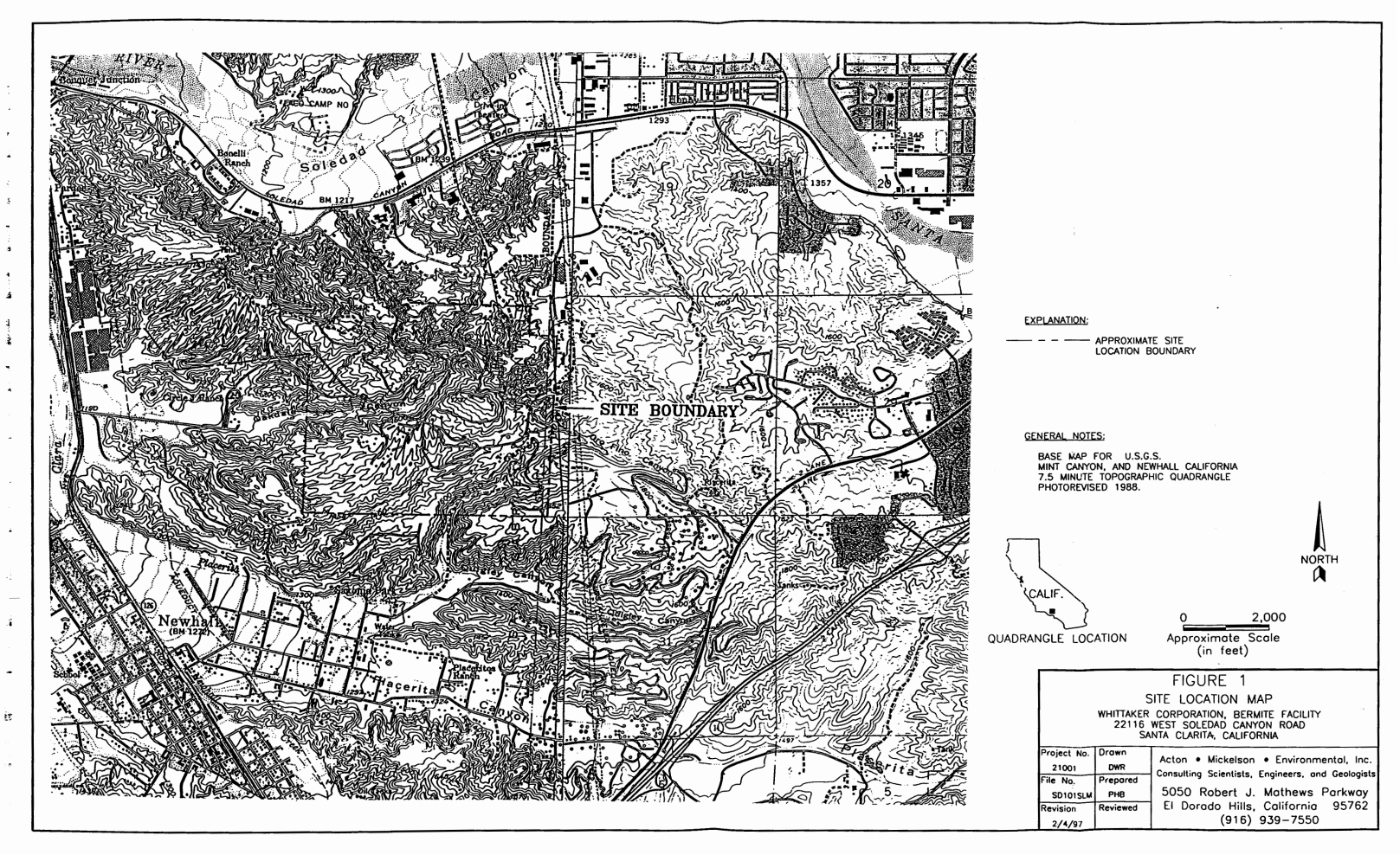
TABLE 3
BACKGROUND WATER QUALITY PARAMETERS

Well No.	Date Sampled	Gross Alpha (pCi/l)	Gross Beta (pCi/l)	Lead (mg/l)	Fluoride (mg/l)	Nitrate (mg/l)	Turbidity (NTUs)
Detection I	Limits			-0.01* 0.0002	0.1	0.4	0.2
MW-1	10/04/88	0.4 ± 2	0.7 ± 2	< 0.01	b		
	01/27/93	0 ± 1	4 ± 2	< 0.01	0.2		
	06/09/93	0.4 ± 1	0.7 ± 2	< 0.01	0.2	3.9	0.4
	07/14/93	2 ± 2	0 ± 2	< 0.01	0.4	4.8	0.9
	08/11/93	1 ± 1	4 ± 4	< 0.01	0.3	4.8	0.9
	09/22/93						0.5
	03/10/94					ND	
	06/22/94	2 ± 2	4 ± 2	< 0.0002	0.2	3.6	1.0
MW-3	10/04/88	0.7 ± 1	2 ± 3	< 0.01			
	01/27/93	0.8 ± 1	2 ± 2	< 0.01	0.3		
	06/09/93	2 ± 1	1 ± 2	< 0.01	0.2	1.6	<0.2
	07/14/93	2 ± 2	1 ± 2	< 0.01	0.3	2.1	<0.2
	08/11/93	4 ± 2	3 ± 4	<0.01	0.2	2.2	0.3
	09/22/93						<0.2
	03/10/94					1.4	
	06/22/94	1.0 ± 1	2 ± 2	4.9	0.2	3.6	0.3
MW-5°	06/22/94	1.0 ± 1	3 ± 2	< 0.0002	0.2	3.6	0.9
MW-6°	06/22/94	0.1 ± 1	2 ± 2	< 0.0002	0.2	3.8	0.8
MW-10°	06/22/94	0.4 ± 1	4 ± 2	< 0.0002	0.2	3.7	0.8

^{*}Detection limit lowered from 0.01 to 0.0002 mg/l on 6/22/94.

^bSample was not taken.

Samples collected from monitoring wells MW-5, MW-6, and MW-7 during the twenty-third sampling event were analyzed for the background water quality parameters because of a repeated tolerance interval exceedence for sodium during previous sampling events.



!		

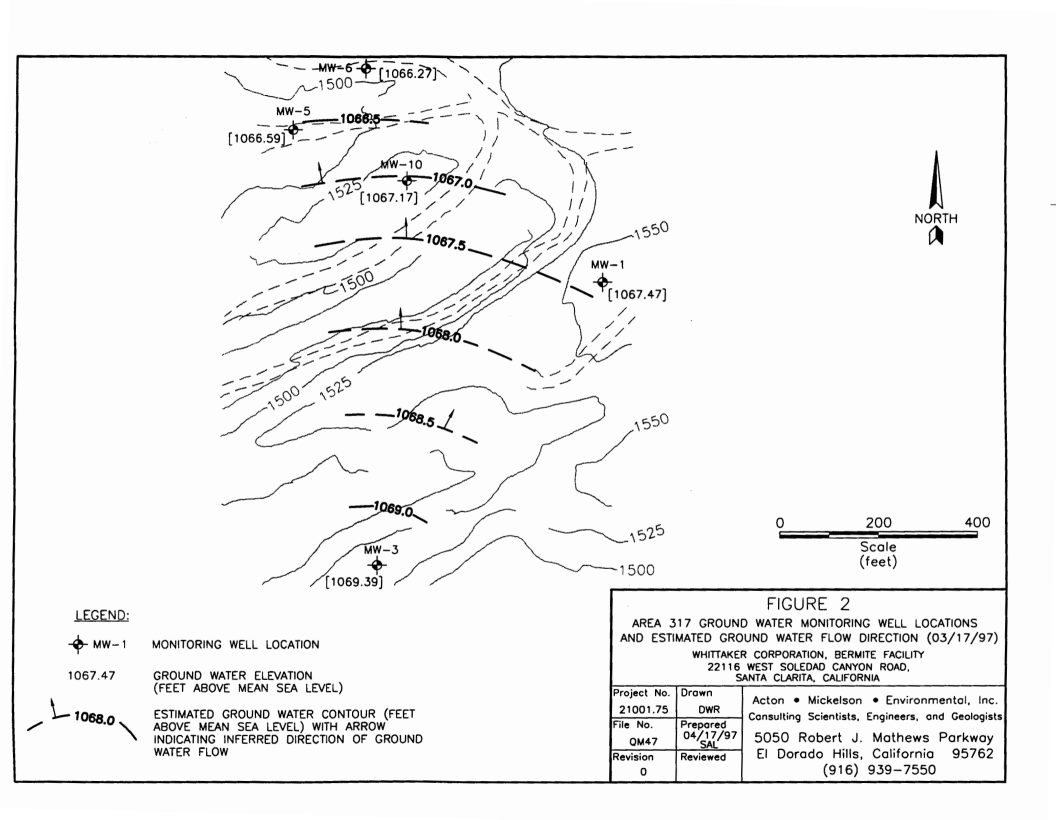
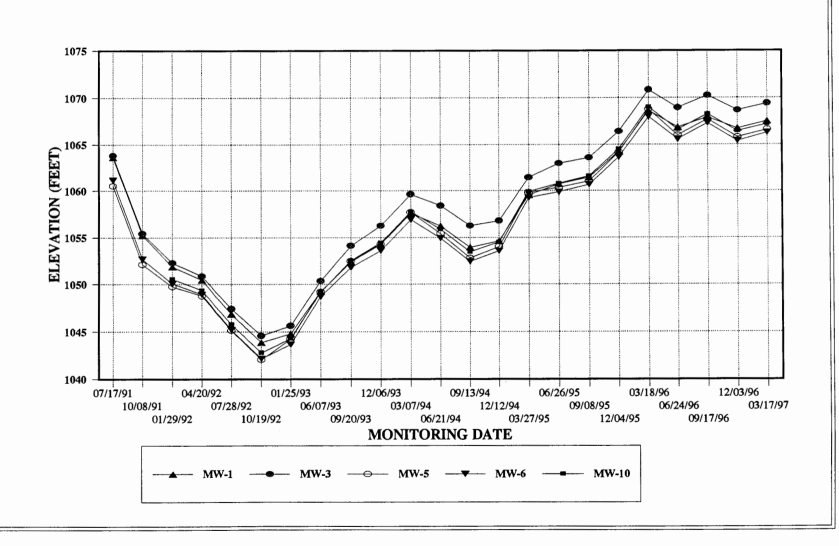


FIGURE 3

RCRA GROUND WATER MONITORING WELLS POTENTIOMETRIC SURFACE ELEVATIONS



APPENDIX A DOCUMENT SUBMITTAL CHRONOLOGY

APPENDIX A

DOCUMENT SUBMITTAL CHRONOLOGY

The following documents have been submitted to Cal-EPA and U.S. EPA, Region IX, in fulfillment of the Closure Plan regarding ground water monitoring at Areas 317 and 342:

- Whittaker Corporation, Bermite Division, Santa Clarita, CA CAD064573108, Facility Closure Plan Modifications, April 1987.
- Revised Ground Water Monitoring Plan for the 317/342 Area, October 8, 1987.
- Proposed Interim Status Ground Water Monitoring Sampling and Analysis Program, December 1987.
- Documentation Report--Construction and Development of Wells for Ground Water Monitoring of the 342 and 317 Areas, February 1988.
- Verification Sampling Results at Selected RCRA Units, March 1988.
- RCRA Ground Water Monitoring System--Proposed Final Configuration, May 1988.
- Ground Water Sampling and Analysis Plan, August 1988.
- RCRA Ground Water Sampling, Quarterly Sampling Report No. 1, December 1988.
- RCRA Ground Water Sampling, Quarterly Sampling Report No. 2, March 1989.
- RCRA Ground Water Sampling, Quarterly Sampling Report No. 3, July 1989.
- Specific Plan for a Ground Water Quality Assessment Program, June 1989.
- Interim Response Action Plan, 317 Area Soil and Ground Water Remediation, June 1989.
- Site Ground Water Sampling and Analysis Plan, Appendix IV of 40 CFR 264.
- RCRA Ground Water Sampling, Quarterly Sampling Report No. 4, September 1989.
- Statistical Analysis--Well MW-2 Versus MW-1 and MW-3, October 1989.
- RCRA Ground Water Sampling, Quarterly Sampling Report No. 5, March 1990.
- RCRA Ground Water Sampling, Quarterly Sampling Report No. 6, May 1990.
- RCRA Ground Water Sampling, Quarterly Sampling Report No. 7, June 1990.
- RCRA Ground Water Sampling, Quarterly Sampling Report No. 8, October 1990.
- RCRA Ground Water Sampling, Quarterly Sampling Report No. 9, January 1991.
- RCRA Ground Water Sampling, Quarterly Sampling Report No. 10, April 1991.

- RCRA Ground Water Sampling, Quarterly Sampling Report No. 11, July 1991.
- RCRA Ground Water Sampling, Quarterly Sampling Report No. 12, October 1991.
- Specific Plan for a Ground Water Quality Assessment Program for the 317 Surface Impoundment Area.
- RCRA Ground Water Sampling, Quarterly Sampling Report No. 13, January 1992.
- Area 317 RCRA Quarterly Ground Water Quality Monitoring Report No. 14 and Report of Monitoring Well MW-10 Installation, January through March 1992.
- Area 317 RCRA Quarterly Ground Water Quality Monitoring Report No. 15, April through June 1992.
- Area 317 RCRA Quarterly Ground Water Quality Monitoring Report No. 16, July through September 1992.
- Water Quality Monitoring and Response Plan for the Interim Status Area 317 Surface Impoundment, October 1992.
- Area 317 RCRA Quarterly Ground Water Quality Monitoring Report No. 17, October through December 1992.
- Area 317 RCRA Quarterly Ground Water Monitoring Report No. 18, January through March 1993.
- Area 317 RCRA Quarterly Ground Water Monitoring Report No. 19, April through June 1993.
- Area 317 RCRA Quarterly Ground Water Monitoring Report No. 20, July through September 1993.
- Area 317 RCRA Quarterly Ground Water Monitoring Report No. 21, October through December 1993.
- Area 317 RCRA Quarterly Ground Water Monitoring Report No. 22, January through March 1994.
- Area 317 RCRA Quarterly Ground Water Monitoring Report No. 23, April through June 1994.
- Area 317 RCRA Quarterly Ground Water Monitoring Report No. 24, June through September 1994.
- Area 317 RCRA Quarterly Ground Water Monitoring Report No. 25, October through December 1994.
- Area 317 RCRA Quarterly Ground Water Monitoring Report No. 26, January through March 1995.

TABLE B-2 WELL STABILIZATION TESTS WHITTAKER CORPORATION, BERMITE FACILITY

Well	Time and Date	Temperature ("C.)	pΗ	Specific Conductance (µmhos)*/cm²	Turbidity (NTUs) ⁵
MW-1	1120 (03-18-97)	23.4	7.97	768	2.91
	1515 (03-18-97)	23.4	7.96	799	3.15
	0720 (03-19-97)	22.1	8.10	781	1.93
MW-3	1125 (03-18-97)	24.2	7.94	592	6.55
	1520 (03-18-97)	24.5	7.94	606	1.54
	0725 (03-19-97)	23.8	8.02	613	1.36
MW-5	1110 (03-18-97)	23.0	7.95	544	3.17
MI W-3	1505 (03-18-97)	23.0	7.91	555	2.60
	0710 (03-19-97)	22.1	8.19	552	1.76
MW-6	1105 (03-18-97)	23.2	7.73	587	14.87
141 44 -0	1500 (03-18-97)	23.9	7.94	582	4.28
	0705 (03-19-97)	22.7	8.27	588	2.74
MW-10	1115 (03-18-97)	22.9	7.75	628	7.08
141 44 -10	1510 (03-18-97)	23.1	7.90	628	4.18
	0715 (03-19-97)	22.3	8.06	631	2.52

^aμmhos - micromhos.
^bNTUs - nephelometric turbidity units.

TABLE B-3

LABORATORY ANALYTICAL METHODS AND SAMPLE VOLUME AND CONTAINER REQUIREMENTS AREA 317 GROUND WATER MONITORING WELLS WHITTAKER CORPORATION, BERMITE FACILITY

Constituent	Analytical Method	Sample Volume	Container Type
Ground Water Monitoring Parameters pH/Specific Conductance Total Organic Carbon Total Organic Halogen Trichloroethylene Sulfate/Chloride Sodium/Iron/Manganese	4500-HB/2510B EPA 415.1 EPA 9020 EPA 624 4110B 200.7/200.7/200.8	16 oz. 250 ml. 250 ml. 3 x 40 ml. 32 oz. 32 oz.	Plastic Amber glass-TFE cap Amber glass-TFE cap Amber glass-TFE cap Plastic Plastic

APPENDIX C CHAIN-OF-CUSTODY FORMS



								QI	JARTI	RLY				abo.	rator	y Copy	(1.of	3)		
Client : Bermite Division o	of Whittaker		Τ	Τ		TES	T DESC	RIPTIO	NS - S	ee rev	erse s	ide for C	Container,	Pres	ervativ	e and Sar	mpling infor	mation		
Address: 22116 W. Soledad C	Canyon Road																		T	
Saugus Phone : (805)259-2242 Fax : (805)259-2244	CA 91350	2-4570		RSE SIDE**						<u>\(\C \)</u>										Sealed?
Project Name : MW1 Qtrly Contact person : Mr. Glen A Purchase order Number: QA/QC report required: Yes	lbun-nur	oring	e(C) Grab(G)	**SEE REVERSE	ble			N 3.3 A		NOT										Cond.:
Sampler(s):Comp sampler setup Date:	/ / Time:	<u>a</u>	Composite(C)		(NP) Non-Potabl	×	H2504 A	250ml (at)-4250g HN 3.3	∢	ABC	Ma, wa									Sample
Rush Results due By:/_		·—·—		a	N (AN)	E.C., pH 1602.(p)	250ml(at)-H ₂ SO ₄	nl(at)-	. cl, so, 32oz.(p)	40ml (v) 7	2.(p)-H			1) . (:0
Date Received:/_/_	2021	•	of Sampling	of Sample	Potable		ļ		t Chem - Cl, Cont. 320		Trace Metals-P - Fe, Mn, Na Cont. 32oz.(p)-HNO3 A	•		P	n					Temp
amp Location/Description		Time Sampled	Type	Type	(P)	Wet Chem Cont.	TOC Cont.	TOX Cont.	Wet	EPA 624 Cont.	Trac									Sample
1 MW1/A/34	3/19/97	900	G	MW		1														
2 MW1/B/34	M	0902	G	MW			1													
3 MW1/C/ 34	11	0704	G	MW				1												
4 MW1/H/34	11	0906		MW					1										Ţ	
5 MW1/0/34	U	0908	G	MW						3									T	
6 MW1/R/34	V	0107	G	MW							1									
												·				···			-	
fisc. Notes:			Rel	inqu	ishe	g sy:	3-19	Date:	7.4	e: R	elinqu	ished By:	3/	ate:	Time:	Relin	quished By:	Da	ate:	Time:
· · · · · · · · · · · · · · · · · · ·	Returned to Clic	ent/	Rec	eive	d by	1		Pate:	Tim	e: R	eceive	a pa:	\mathcal{L}) ate:	Time 1919		ved by	Da	ate:	Time:
ornorate Offices & Laboratory				- T	-			Office				- 14 1 Y	71	- //				= 111		

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Visalia, California

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Mobile: (209) 737-2399



Laboratory Copy (1 of 3) **OUARTERI Y**

Clie	nt : Bermite Division of Whitt	aker		Γ	T	Ī	TES	T DESC	RIPTIO	NS - S	ee rev	erse s	ide for Co	ntainer, F	reservativ	e and Samp	oling inform	nation			
Addr	ess: 22116 W. Soledad Canyon R	oad		İ											Ţ ,						
Phone Fax	Saugus CA e: (805)259-2242 : (805)259-2244	91350	-4570		REVERSE SIDE**						S/H									Seal ed?	
Conta Purch	ect Name : MW3 Qtrly 317 Are act person : Mr. Glen Abun-nur mase order Number: creport required: Yes		ring	ite(C) Grab(G)	**SEE REV	table		₹	HIG3 A		NO	₽. ¥						6		le Cond.:	- 1
Comp Rush	sampler setup Date://_ Results due By:/_/_ Received:/_/_	_ Time:	_:_	Sampling: Composite(C)	Sample	le (NP) Non-Potabl	1602.(p)	250ml(at)-H ₂ SO ₄	250ml(at)-H ₂ 80 ₄	- Cl, SO ₄ 3202.(p) A	40ml(v) ABC	Metals-P - Fe, Mn, t. 32oz.(p)-HNO3 /								Temp °C: Sampl	1
Lab N Samp Num	Location/Description		Time Sampled	ype of	Type of S	(P) Potable	Wet Chem Cont.	TOC Cont.	TOX Cont.	Wet Chem Cont.	EPA 624 Cont.	Trace Met Cont.								Sample Te	
1 M	W3/A/34		0920	_	MW		1												$oxed{igspace}$		١
2 M	W3/B/3¶		922	G	MW			1]
3 M	N3/C/34		6924	G	MW				1												
4 M	M3/H/34	11	926	G	MW					1											1
5 M	M3/0/ 3H	11	3929	G	MW						3									\perp	
6 M	43/R/31/j	16	0927	G	M.							1									1
																					1
	Notes:			Re	linqu	ish		3.1	Date:	7in	ne: R	telinqu	uished By:	- 3/10a	te: /e Time:	Relino	uished By:	Da	ite: 1	Time:	
Lab D	Sample Dispostion: isposal:// Returned of Disposal: Date Ret		ent _/	Re	eive	d by	3	. 3/	Date:	Ogy	re: R	eceive B	ed by:	WOO	311419	Receiv	ed by	Da	ite: 1	Time:	

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Clie	nt : Bermite Division of White	taker		T	Τ	T	TES	T DESC	RIPTIO	NS - S	ee rev	verse s	ide for Co	ntainer, F	reservati	ve and Samp	oling inform	nation		
Addr	ess: 22116 W. Soledad Canyon F	Road														T				
Phon Fax	Saugus CA e : (805)259-2242 : (805)259-2244	91350	2-4570		REVERSE SIDE**						8									Sealed?
Conta Purci QA/Qd Samp Comp	ect Name : MW5 Qtrly 317 Are act person : Mr. Glen Abun-nur nase order Number: C report required: Yes Ler(s): Sampler setup Date:// Results due By:/_/	No	<u> </u>	ing: Composite(C) Grab(G)	**SEE	(NP) Non-Potable	16oz.(p) Д	250ml(at)-H ₂ SO ₄ A	250ml(at)-4g60r HN03 A-	32oz.(p) A	40M CA BY NO H	Fe, 7 (p)-HNC		907	3/19					°C: Sample Cond.:
	Received:/_/	2 2 Date	Time	Type of Sampling:	Type of Sample	(P) Potable	Wet Chem - E. Cont. 160	TOC Cont. 250	TOX Cont. 250	Wet Chem - Cl Cont. 320	EPA 624 Cont. 40m	Trace Metals-P Cont. 32oz								Sample Temp°
Num	A south to the second s	1 '	Sampled		┼	5		٢	70	3	<u> </u>	F							<u> </u>	Š
	N5/A/34	3/19/97	370	G	MW		1					<u> </u>							ļ <u>.</u>	4-4
-+	M2/B/ 34	11	0822	G	┼			1											<u> </u>	44
	W5/C/34		0874	G	MW				1											44
4 M	W5/H/3·f		0877	!	MW					1		L								
5 M	N5/0/34	11	683-8	G	MW						3									
6 M	\ 5/R/3 √	N	0827	G	MW							1								
	Notes:			Re	l inqu	uishe	d Byo			1.45		Relinqu	uished By:	3/1/2	te: Time	: Reling	uished By:	Da	te: T	ime:
Lab D	Sample Dispostion: isposal:// Returned of Disposal: Date Ret	d to Clie t/_	ent _/	Re	ceive	d by	De.	- 3	Pate:	Tim	e:	Receive B	od by:	Dat			ed by:	Da	te: T	ime:

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Client : Bermite Division of Whittaker		T		TES	T DESC	RIPTIO	vs - s	ee rev	erse s	ide for Co	ntainer, P	reservativ	and Samp	ling inform	nation		
Address: 22116 W. Soledad Canyon Road	İ															T	
Saugus CA 91350 Phone : (805)259-2242 Fax : (805)259-2244 2-4570		REVERSE SIDE**						\$\frac{1}{S}									Sealed?
Project Name : MW6 Qtrly 317 Area Monitoring Contact person : Mr. Glen Abun-nur Purchase order Number: QA/QC report required: Yes No	te(C) Grab(G)	*SEE	ble			HNO 3		S			, 0	3/0					Sample Cond.:
Sampler(s):	Composit		Non-Potab		2504	250ml(at)- UgSO_t /		T (X)	Mn, Na O3								Sample
Comp sampler setup Date:// Time::_				H ₍₀₎ :	250ml(at)-H ₂ SO4	£	33		Fe,								
Rush Results due By://	Ē	e e	(NP)	, . Z	l es	ml(a	. Cl, SO4 32oz.(p)	40ml (v)	P - 2								ູ່:
Date Received:/_/ Lab Number:	of Sampling:	6	Potable	Chem - E.C. ont. 160z.			•		Hetals-P - Fe, Mn, ht. 32oz.(p)-HNO3								Temp
Samp Location/Description Date Time Num Sampled Sample		Type	(e)	Wet Che Cont.	TOC Cont.	TOX Cont.	Wet Chem Cont.	EPA 624 Cont.	Trace Me Cont.								Sample
1 MH6/A/34 3/19/97 0800	G	MW		1													
2 MH6/B/34j 11 880	G	MW			1												
3 MH6/C/34 il 0804	G	MW				1											
4 MW6/H/34 (1 980)	G	MW					1										
5 MH6/0/34 11 0805	G	MN						3									
6 MW6/R/34 11 088	G	MW							1								
															- No. 32	-	
Misc. Notes:	Re	l inqu	rishe	d By:	3- <i>/</i> /	Date:	Tin 9'.'5		el inqu	uished by:	3 pat	e: Jime:	Relinqu	uished By:	D	ate:	Time:
Final Sample Dispostion: Lab Disposal:// Returned to Client Meth. of Disposal: Date Ret//	Re	ceive	d by	2	· 3	Pate:			3	DILOY	Dat	3/1990	Receive	ed by:		ate:	Time:

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Client : Bermite Division of Whittaker	Τ	Π	Π	TES	T DESC	RIPTIO	NS - S	ee rev	verse s	ide for Co	ntainer,	Preserva	tive a	ınd Samp	oling infor	mation			
Address: 22116 W. Soledad Canyon Road																			¥
Saugus CA 91350 Phone : (805)259-2242 Fax : (805)259-2244 2-4570	G	REVERSE SIDE**																	Sealed?
Project Name : MW10 Qtrly 317 Area Monitoring Contact person : Mr. Glen Abun-nur Purchase order Number: QA/QC report required: Yes No Sampler(s) Comp sampler setup Date:/ Time:: Rush Results due By:/ Date Received:/ Lab Number:		ole **SEE	Potable (NP) Non-Potable	hem - E.C., pH nt. 16oz.(p)	nt. 250ml(at)-H ₂ SO ₄	nt. 250ml(at)- 4269 4 Hrv 3	them - Cl, SO, nt. 32oz.(p)	12. 40m(cv) NO H/S	e Metals-P - Fe, Mn, Na nt. 32oz.(p)-HNO3			497		611					le Temp °C: Sample Cond.:
Samp Location/Description Date Time Num Sampled	7,78	Type	(9)	Wet Chem Cont.	TOC Cont.	TOX Cont.	Wet Chem Cont.	EPA 624 Cont.	Trace Mc Cont.								į		Sample
1 MH10/A/34 3/19/97 0840	G	MW		1															
2 MW10/B/34 11 0842	G	MW			1														
3 MW10/C/34 11 0844	G	MW				1													
4 MN10/H/34 11 0846	G	MW					1												
5 MW10/0/34 11 0849	G	MW						3]	
6 MW10/R/ 34 11 8847	G	MU							1										
Misc. Notes:	Rel	inqu	rish	stay:	3-19	Date:	Tin T. 4	ne:	Relinqu	uished 89?	- 3/19/	ite: T	ime:	Relino	uished By:		Date:	Tim	æ:
Final Sample Dispostion: Lab Disposal:/ Returned to Client Meth. of Disposal: Date Ret//	Rec	eive	d by	- 3	-	Date:			3B	ed by:		143/1	919	Receiv	ved by:	_	Date:	Tim	e:

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Mobile: (209) 737-2399



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Cli	ent : Bermite Division of White	taker		1	Π	Γ	TES	T DESC	RIPTIO	IS - See r	everse sid	le for Cont	ainer, Pre	servative	and Sampli	ing informa	tion		
Add	ress: 22116 W. Soledad Canyon R	Road													T]
Saugus CA 91350 Phone : (805)259-2242 Fax : (805)259-2244 2-4570									H15										Seal ed?
Project Name : MW5-1A Qtrly 317 Area Monitor.						Non-Potable	4	# 502 H	🛩			2	6//						ole Cond.:
	oler(s):	Time:	<u> </u>	soduo		9-re	H2504	# 03 2#	AB			Y	(7)						Sampl
Rush Results due By:/						(NP)	250ml(at)-H ₂ SO ₄	250ml(at)-H 250g 析いの3	(^) #0#										Temp °C:
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Corporate Offices & Laboratory P.O. Box 272 / 853 Corporation Street Santa Paula. CA 93061-0272

TEL - 905- 650,0010

Office and Laboratory 2500 Stagecoach Road Stockton, CA 95215 TEL: (209) 942-0181

APR - 4 1997

Field Office Visalia, California TEL: (209) 734-9473 Mobile: (209) 737-2399



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Laboratory Copy (1 of 3)

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Corporate Offices & Laboratory
P.O. Box 272 : 853 Corporation Street
Santa Paula. CA 93061-0272
TEL: (805) 659-0910

Office and Laboratory 2500 Stagecoach Road Stockton. CA 95215 TEL: (209) 942-0181

APR - 4 1997

Field Office Visalia, California TEL: (209) 734-9473 Mobile: (209) 737-2399

APPENDIX D SAMPLE ANALYSES REQUEST FORMS

SAMPLE ANALYSIS REQUEST

Sampling Inform	ation				3411	. 2			
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				Analy	sis !	Requi	red		
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SAMPLE ANALYSIS REQUEST

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Date Samples Re	ceived: 3/1	9/97					······································	
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			Analy	ysis 1	Requi	red		
		P.K.	Tac	Tox	SOCFATE, CHLORNOE	EPA 624 TCE ONLY	TRUN, MANGANESE, SODIUM	
Sample I.D.	Laboratory I.	.D.						
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MW5/B/34		2	X					
MU5/C/34		3		X				
MW5/H/34	-1	4			4			
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mw5/R/3H	4 -1	٥					X	
mw6/A/34	707025	5-1 ×						
MW6/B/34		-2	*					
mwb/c/34		-3		X		-		
mw6/H/34		-4		-	X			-
mw6/0/34		-5				X		
	_	-10					×	

SAMPLE ANALYSIS REQUEST

Sampling Inform	ation			O.	2	15		
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Sampler Name: G	LENABOW-NUR/TIM BRICK				-)259	1-224	1
Name of Person	Receiving Samples:	Sta	cey	Bei	crit	19±	Ch_	
Date Samples Re	ceived: 3/19/0	17	/					
Internal Temperature of Sample Container:								
Notes on Sample	es:							
			Analy	sis I	Requi	red		
		P#, EC	70 C	XOX	SULFATE, CHCANDE	EPA 624 TCE ONLY	IRON, MANGANESE, SODIUM	
Sample I.D.	Laboratory I.D.							
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MU10/C/34	-3			X				
MW10/H/34	-4				X			
mw10/0/34	-5					×		
M418/R/34	4 -6						4	
MW5/8/34/1A	702024-1		*			-		
MW5/C/34/1A	-2			*				
MW5/0/34/1A	-3					X		
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MW6/C/34/1A	1 -2			X				
MW6/0/34/1A	√ -3					X		

APPENDIX E FGL QUALITY ASSURANCE/QUALITY CONTROL PROGRAM

FGL ENVIRONMENTAL

QUALITY ASSURANCE MANUAL

Reviewed by	fate	6/4/96
•	Yee Ren, M.S.	Date
	Data Validator	
Concurred by	Jun Jun	4/4/96
•	Kurt Wilkinson, B.S.	Date /
	Quality Assurance Director	
Approved by	Canul H. Nelson	6/4/96
	Darrell H. Nelson, B.S.	Date
	Laboratory Director	

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2 Table of Contents	2.5	5-22-96	3
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4 Laboratory Organization and Responsibilities	2.4	5-22-96	18
5 Quality Assurance Objectives for Measurement Data in Terror of Precision, Accuracy, Completeness, and Detection Limits for Reporting	ms 2.2	5-22-96	33
6 Sampling Procedures	2.3	5-22-96	16
7 Sample Custody	2.1	10-3-94	7
8 Calibration Procedures and Frequency	2.3	5-22-96	6
9 Analytical Procedures	2.3	5-22-96	14
10 Data Reduction, Validation, and Reporting	2.2	5-9-95	2
11 Internal Quality Control Checks	2.2	5-22-96	19
12 Performance and System Audits	2.3	5-22-96	31
13 Preventative Maintenance	2.2	5-9-95	5
14 Specific Routine Procedures Used to Assess Data Precision, Accuracy & Completeness	2.2	5-9-95	3
15 Corrective Actions	2.1	10-3-94	3
16 Quality Assurance Reports to Management	2.1	10-3-94	3
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Date: May 22, 1996

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- Table 5-4 Quality Assurance Objectives for Ambient Air / Air Pollution Methods
- Table 6-1 Recommended Sample Containers, Preservation and Holding Times
- Table 8-1 GC/MS Volatile Organic and Air Toxics Key Ion Abundance Tuning Criteria using BFB
- Table 8-2 GC/MS Semivolatile Organic Key Ion Abundance Tuning Criteria using DFTPP
- Table 9-1 Specific Analytical Drinking Water Methods
- Table 9-2 Specific Analytical Wastewater / Hazardous Waste Liquid Methods
- Table 9-3 Specific Analytical Solid Waste / Hazardous Waste Methods
- Table 9-4 Specific Analytical Ambient Air / Air Pollution Methods
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- Figure 15-1 Corrective Action Report Form
- Figure 16-1 FGL QC Inspection Summary Report Form

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QA Plan Description

FGL Environmental's Quality Assurance Policy

"Management and staff are committed to providing data that is scientifically valid, legally defensible and of known precision and accuracy in order to meet or exceed the data quality objectives of our clientele."

"Management and staff are committed to providing our clientele with services that satisfy their definitions and expectations of quality."

This manual describes the protocols utilized by FGL Environmental to ensure adherence with this policy. It outlines quality control procedures to be used with field and analytical methods. It also includes the individual analysis data quality objectives. Detailed project-specific FGL standard operating procedures to supplement this manual are provided when requested or deemed necessary.

FGL's QA manual is based on the 16 essential elements contained in the U.S. Environmental Protection Agency manual "Interim Guidelines and Specifications for Preparing Quality Assurance Project Plans," QAMS-005/80. Two additional appendices have been added. The first appendix lists the laboratory equipment. The second describes FGL's document control program.

As of the date of revision, this manual reflects the current quality assurance program in effect and procedures in the process of being adopted.

References for this manual include field and laboratory methods published by the U.S. Environmental Protection Agency and other agencies mainly through the following sources:

- (1) "Standard Methods for the Analysis of Water and Wastewater," 18th Edition, 1992.
- (2) "Methods for Chemical Analysis in Waters and Waste," (MCAWW) EPA-600/4-79-020
- (3) "Methods for the Determination of Organic Compounds in Drinking Water," EPA Method Book, EPA-600/4-88-039, December 1988.
- (4) "Methods for the Determination of Organic Compounds in Drinking Water-Supplement I," EPA Method Book, EPA-600/4-90- 020, July 1990.
- (5) "Methods for the Determination of Organic Compounds in Drinking Water-Supplement II," EPA Method Book, EPA-600/4- 90-020, July 1990.
- (6) "Methods for Organic Chemical Analysis of Municipal and Industrial Wastewater," EPA Method Book, EPA 600/4-82-057, July 1982.
- (7) "Methods for Evaluating Solid Waste," EPA Method Book, SW-846, 3rd edition, and Proposed Revisions
- (8) "Prescribed Procedures for Measurement of Radioactivity in Drinking Water," EPA Method Book, EPA-600/4-80-032, August 1980.
- (9) "Handbook for Sampling and Sample Preservation of Water and Wastewater," EPAMethod Book, EPA-600/4-82-029, September 1982.
- (10) "Eastern Environmental Radiation Facility Radiochemistry Procedures Manual," EPA Method Book, EPA 520/5-84-006, August 1984.

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QA Plan Description

References continued

- (11) "Environmental Measurements Laboratory Procedures," HASL-300, 27th Edition, February 1992.
- (12) "Compendium of Methods for the Determination of Toxic Organic Compounds in Ambient Air," EPA Method Book, EPA-600/4-, June 1988.
- (13) "Technical Assistance Document for Sampling and Analysis of Toxic Organic Compounds in Ambient Air," EMSL, EPA Method Book, EPA-600/4-83-027, 1983

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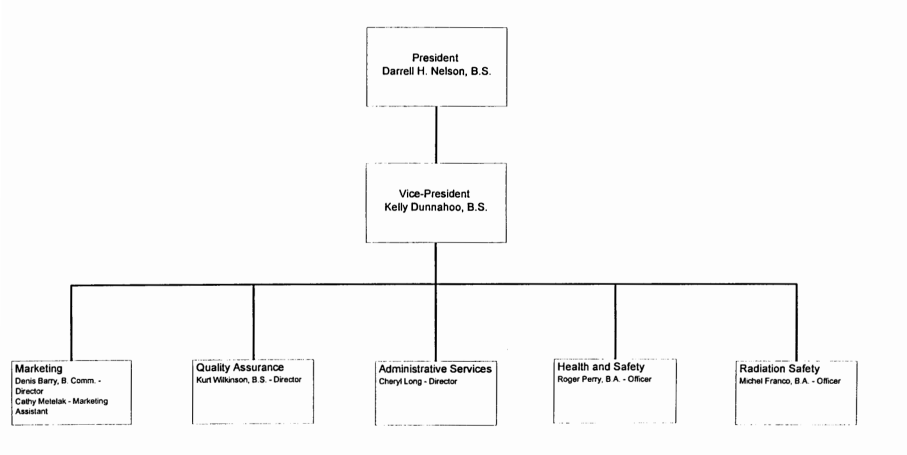
Laboratory Organization and Responsibilities

Personnel of FGL Incorporated

Our commitment to providing a superior service requires the high caliber staff employed by FGL. These professionals have considerable experience to meet the technical demand of environmental analysis. Figures 4-1 - 4-4 show the organization for the corporation and each laboratory or field service facility. Following the figures are qualification summaries for key personnel.

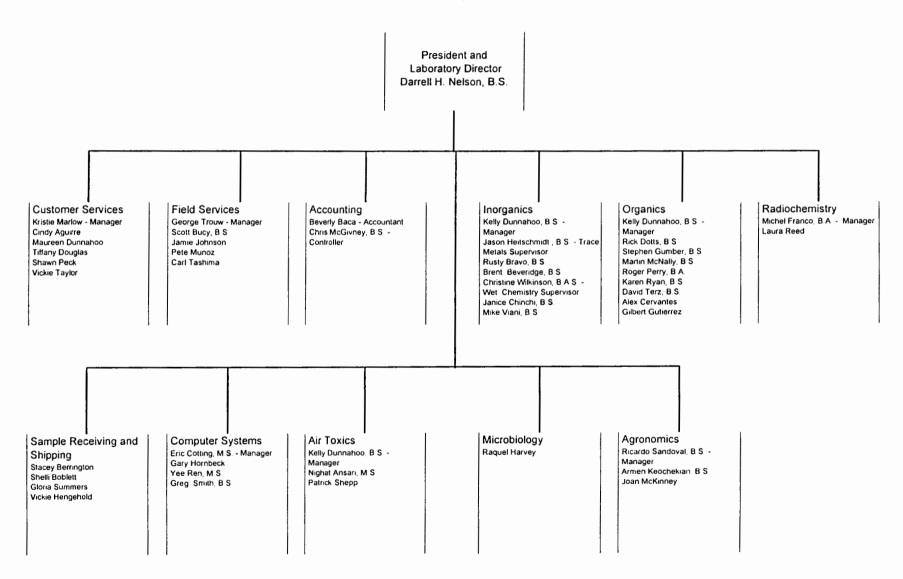
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Laboratory Organization and Responsibilities Figure 4-1 Organization Chart Corporate Personnel



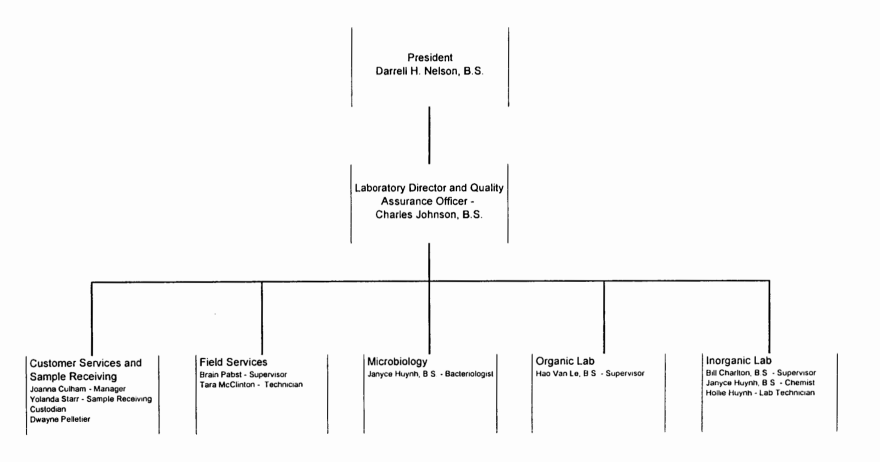
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Laboratory Organization and Responsibilities Figure 4-2 Organization Chart Santa Paula Laboratory



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Laboratory Organization and Responsibilities Figure 4-3 Organization Chart Stockton Laboratory



* * *

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Laboratory Organization and Responsibilities Figure 4-4 Organization Chart Visalia Field Office

President
Darrell H. Nelson, B.S.

Field Services
Neil Jessup, B.S.

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Laboratory Organization and Responsibilities

Key Personnel Qualification Summary - Santa Paula

Darrell H. Nelson

Current Responsibilities:

President and Lab Director

- To oversee all operational aspects of the Santa Paula and Stockton laboratories
- To guide and direct managers towards the corporation's goals
- To report to the Board of Directors and the stockholders

Work Experience:

Assistant Manager, FGL

- Chemical analysis of drinking and waste waters, soils and plant materials
- Supervision of a number of lab operations including work scheduling and field services
- Customer Interface

Formal Education:

- B.S. (1970) in Soil and Water Science, University of California, Davis.

Continuing Education:

- University of Southern California (USC) School of Business Administration, "Customer Service Management," April 1990.
- Hazardous Waste Operations and Emergency Response Training, OSHA 29 CFR 1910.120, 40 hr plus annual refresher.
- American Chemical Society (ACS), "Quality Assurance for Analytical Chemistry," Nov. 1991.
- University of California Davis (UCD), "Marketing Professional Services," Feb. 1988.
 University of California Davis (UCD), "Guerrilla Marketing," Feb. 1988.
 University of California Davis (UCD), "Enhancing Sales Skills," Feb. 1988.

- American Water Works Association, "Approved Water Sampling Procedures," March
- California Agricultural Leadership Program, 1976 1978.
- NPDES Requirements for Industrial and Construction Site Storm Water Discharges, ASCE, Feb. 1992.
- Senate bill #198, compliance training (State Fund Insurance)
- Nevada Nuclear Associates, "Fundamentals of Radiochemistry", February 1994.

Memberships:

Professional:

- American Chemical Society
- American Water Works Association
- Association of California Testing Laboratories

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Laboratory Organization and Responsibilities

Key Personnel Qualification Summary - Santa Paula

Kelly Dunnahoo

Current Responsibilities:

Vice President

- To support the president in all operational, technical and strategic aspects relating to the corporation.

Inorganic, Organic and Air Toxics Lab Manager:

- Organization of work flow in the above departments
- Personnel training, methods development, instrument troubleshooting
- Response to specific customer enquiries

Work Experience:

- 10 years experience in environmental and geochemical analysis
- 6 years of supervisory and management roles in organic laboratory operations

Formal Education:

- B.S. (1987) in Biochemistry, University of California, Los Angeles

Continuing Education:

- AOAC, "QA for Analytical Labs", November 1991.

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Laboratory Organization and Responsibilities

Key Personnel Qualification Summary - Santa Paula

Denis Barry

Current Responsibilities:

Marketing Director

- Responsible for marketing of FGL's services to city, county and state and federal agencies in addition to private companies
- Compilation and implementation of FGL's marketing plans
- Responsible for FGL's informational and promotional materials

Work Experience:

- 6 years experience in marketing analytical services
- Marketing consultancy experience focused on small to medium sized companies
- International programs targeted mainly at the European Economic Community for small U.S. companies

Formal Education:

- B. Comm (Bachelor of Commerce), University College, Dublin, Ireland

Continuing Education:

- Computer Appreciation - Moorpark College, 1989

Memberships:

- Vice President - Irish American Club of Ventura County

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Laboratory Organization and Responsibilities

Key Personnel Qualification Summary - Santa Paula

Cheryl Long

Current Responsibilities:

Administrative Services Director

- Management of Santa Paula Office
- Organize and prioritize work load
- Document Control

Work Experience:

20 years' management experience environmental field

- Supervision of office personnel, customer service, marketing, data input, invoicing, accounts receivable, bacteriologist, laboratory technician, sample receiving, and field sampling.
- Additional experience in analyzing animal feed rations/rumen digestions.
- Environmental laboratory experience to include agricultural and air testing
- Physical testing construction and inspection laboratory affidavits, Notary Public

Formal Education:

- Associates of Arts Degree, Porterville College, Porterville, California

Continuing Education:

- Bakersfield College Business Education and Management classes
- Water Treatment Plant Operator I License

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Laboratory Organization and Responsibilities

Key Personnel Qualification Summary - Santa Paula

Kurt Wilkinson

Current Responsibilities:

Quality Assurance Director

- Design, implementation and maintenance of FGL's Quality Assurance Program

Work Experience:

- 10 years experience in environmental testing of drinking water, wastewater, hazardous waste and air analysis
- Additional experience in agricultural testing of soils, plant tissue and food products

Formal Education:

- B.S. (1987) in Biochemistry, California Polytechnic State University, San Luis Obispo

Continuing Education:

- American Chemical Society (ACS), "Environmental Analytical Chemistry, Water and Waste," Nov. 1991.
- American Chemical Society (ACS), "Gas Chromatography/Mass Spectrometry," April 1992.
- Halliburton NUS, "Solving the Mysteries, Collecting Environmental Samples," April 1992.

Memberships:

Professional:

- American Chemical Society (ACS)

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Laboratory Organization and Responsibilities

Key Personnel Qualification Summary - Santa Paula

Roger Perry

Current Responsibilities:

Health and Safety Officer

- Design, implementation and maintenance of FGL's Health and Safety programs
- Organic Department Chemist

Experience:

- 14 years experience as an environmental analytical chemist
 Health and Safety regulations and compliance
- Handling accumulation and disposal of Hazardous Wastes and Radioactive Materials

Formal Education:

- B.A. (1982) in Chemistry, Sonoma State University, Sonoma

Continuing Education:

- Hazardous Waste Operations and Emergency Response Training, OSHA 29 CFR 1910.120, 40 hr plus annual refreshers.

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Laboratory Organization and Responsibilities

Key Personnel Qualification Summary - Santa Paula

Michel M. Franco

Current Responsibilities:

Radiochemistry Lab Manager

- Work scheduling and staff supervision
- Oversee data analysis, data validation and recording
- Method and Standard Operating Procedure development and review for maximizing efficiency and turnaround time for the Radiochemistry Laboratory
- Personnel training and instrument troubleshooting
- Client interface and project management

Radiation Safety Officer

- Conduct personnel training and dosimetry program
- Maintenance of safety systems and spill cleanup kits
- Internal inspections of logs and storage areas
- Perform required leak tests of sealed sources
- Monitor radioactive material receiving, tracking, waste handling and holding areas

Work Experience:

- 8 years experience in radiochemical analysis, general inorganic analysis and organic analysis for TOX, TOC, HPLC and GC.
- Developed instrument analysis experiment for CSUN (1988)
- Sample trouble shooting, processing and department liason at Reference Laboratory (1986)

Formal Education:

- B.A. (1990) Chemistry, California State University, Northridge

Continuing Education:

- Nevada Nuclear Associates, "Fundamentals of Radiochemistry", February 1994.
- Canberra Industries Inc., "Environmental Radioactivity Quantification Workshop", March 1994.
- Health Physics Seminar June 1994.
- Aptec Inc., "Gamma Ray Spectroscopy Interactive Training", January 1995.

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Laboratory Organization and Responsibilities

Key Personnel Qualification Summary - Santa Paula

Jason Heitschmidt

Current Responsibilities

Trace Metals Laboratory Supervisor

- Work scheduling and staff supervision
- Oversee data analysis, data validation and recording
- Method and Standard Operating Procedure development and review for maximizing efficiency and turnaround time for the Radiochemistry Laboratory
- Personnel training and instrument troubleshooting

Work Experience:

- 5 years experience in environmental testing of drinking water, waste water, hazardous waste

Formal Education:

- B. S. (1991) in Zoology, Texas Tech University, Lubbock

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Laboratory Organization and Responsibilities

Key Personnel Qualification Summary - Santa Paula

Christine Wilkinson

Current Responsibilities

Wet Chemistry Laboratory Supervisor

- Work scheduling and staff supervision
- Oversee data analysis, data validation and recording
- Method and Standard Operating Procedure development and review for maximizing efficiency and turnaround time for the Radiochemistry Laboratory
- Personnel training and instrument troubleshooting

Work Experience:

- 6 years experience in environmental testing of drinking water, waste water, hazardous waste
- 3 years supervisory experience in inorganic laboratory operations

Formal Education:

- B. A. & S. (1988) in Biological Sciences and Anthropology, University of California, Davis

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Laboratory Organization and Responsibilities

Key Personnel Qualification Summary - Santa Paula

Ricardo Sandoval

Current Responsibilities:

Agriculture Lab Manager

- Management of Agricultural Department
- Supervision and training of staff

Work Experience:

- 11 years experience in agricultural testing of soils and plant tissue
- Ranch and Nursery experience dealing with irrigation, fertilization and transplantation of avocado trees and horticultural plants

Formal Education:

- B.S. (1985) in Crop Science and Technical Degree in Fruit Science, California Polytechnic State University, San Luis Obispo.

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Laboratory Organization and Responsibilities

Key Personnel Qualification Summary - Santa Paula

Eric Cetting

Current Responsibilities:

Computer Systems Manager

- Design, implementation and maintenance of FGL's Laboratory Information Management System (LIMS)
- Training of personnel in the effective use of LIMS

Work Experience:

- Developed customized LIMS system for FGL
- As research assistant with the University of Wisconsin, was involved in the modification of an existing theoretical computational program designed to model quantum mechanical properties of the Helium Atom

Formal Education:

- B.S. (1981) Chemistry, University of Alaska, Fairbanks
- B.S. (1981) Math, University of Alaska, Fairbanks
- B.S. (1981) Physics, University of Alaska, Fairbanks
- M.S. (1987) Physical Chemistry, University of Wisconsin, Madison

Continuing Education:

- Computer Networking Seminar: Santa Barbara, CA. January 1991
- Nevada Nuclear Associates,"Fundamentals of Radiochemistry", February 1994.

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Laboratory Organization and Responsibilities

Key Personnel Qualification Summary - Stockton

Charles Johnson

Current Responsibilities:

Laboratory Director

- Management of the Stockton Laboratory

- Liason with Santa Paula (Corporate) on Stockton lab's performance, budgets and financial results
- To support the president in all operational, technical and strategic aspects relating to the corporation

Work Experience:

- One year general management with FGL at the Santa Paula laboratory
- One year as Research Assistant University of California, Davis Designing and performing experiments, presentation of results.
- One year as Laboratory Assistant-sample preparation of food crops for pesticide residue analysis.

Formal Education:

- B.S. (1995) - Environmental Toxicology - University of California, Davis

Instrument Experience:

- GC, HPLC, GCMS, and Spectrophotometry

Continuing Education:

- Hazardous Waste Operations and Emergency Response Training, OSHA s9 CFR 1910.120, 40 hours

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Laboratory Organization and Responsibilities

Key Personnel Qualification Summary - San Joaquin

Neil Jessup

Current Responsibilities:

Agronomist/Field service - Visalia

- Technical Representative San Joaquin Valley
- Coordinating sampling and sample pick up in the South San Joaquin Valley
- Consultation with clients on sampling and analysis requirements
- Proposal preparation for contracts on environmental and agricultural testing

Work Experience:

- 12 years experience in areas of field service and sampling
- Considerable background in city, county, state and federal regulations for environmental testing requirements

Formal Education:

- B.S. (1977) in Agronomy, California Polytechnic State University, San Luis Obispo

Continuing Education:

- OSHA 40 hour trained for hazardous waste and emergency response, confined space entry and SCBA
- American Water Works Association, "Approved Water Sampling Procedures," March 1991.
- Halliburton NUS, "Solving the Mysteries, Collecting Environmental Samples," April 1992.

Memberships:

Professional:

- California Agriculture Production Consultants Association (CAPCA)

Civic

- Tulare County Hazardous Waste Advisory Committee

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Quality Assurance Objectives

The quality assurance objectives for accuracy, precision, and Detection Limits for Reporting (DLR) are listed in Tables 5-1 (Drinking Water Methods), 5-2 (Wastewater / Hazardous Waste Liquid Methods) and 5-3 (Solid Waste / Hazardous Waste Solids Methods).

Accuracy - is based on the recovery measurement of a target analyte after known addition to a given sample or representative sample matrix (see section 14.2). Accuracy values are expressed as the percent recovery of the known value, and serve as a reflection of the total measurement error (random and systematic). The acceptance ranges for recovery (%REC-AR) are used for data validation.

Precision - is based on the difference measurement of duplicate data points (see section 14.1). Precision values are expressed as relative percent difference (RPD) and serve as a reflection of the variability in measurement replication. Surrogates are not run in duplicate, therefore RPDs are not applicable. The Maximum Acceptance Value for the RPD's (RPD-MAV) are used for data validation.

Detection Limit for Reporting (DLR) - is the routine detection limit FGL uses for reporting purposes. Method Detection limit studies are performed annually or as specified in the method reference to ensure that the DLR's listed in this section are met. DLR's for surrogates are not listed as surrogates are required for quality control purposes only.

Completeness - FGL is currently introducing controls to document incomplete reports. These are reports that are known to lack information at the time of delivery or reports where we are notified by the client that information is not complete. Future QA manuals will have the results for data completeness documented.

TABLE 5-1 Quality Assurance Objectives for Drinking Water Methods

CONSTITUENT	ACCURACY <u>% REC-AR</u>	PRECISION RPD-MAV	DLR ug/L
EPA Method 501.2			
Bromodichloromethane	70-130	20	0.5
Bromoform	70-130	20	0.5
Chloroform	70-130	20	0.5
Dibromochloromethane	70-130	20	0.5
EPA Method 502.2			
Surrogates			
BFB	70-130	N/A	N/A
Fluorobenzene	70-130	N/A	N/A
Chlorofluorobenzene	70-130	N/A	N/A
Analytes			
Benzene	37-151	30	0.5
Bromobenzene	50-150	30	0.5
Bromochloromethane	50-150	30	0.5
Bromodichloromethane	35-155	30	0.5
Bromoform	45-169	30	0.5
Bromomethane	D-242	30	0.5
n-Butylbenzene	50-150	30	0.5
sec-Butylbenzene	50-150	30	0.5
or but indicate	20-120	50	0.5

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TABLE 5-1 Quality Assurance Objectives for Drinking Water Methods

	ACCURACY	PRECISION	DY D
CONSTITUENT	% REC-AR	RPD-MAV	DLR <u>ug/L</u>
	70 100 111	<u> </u>	<u> </u>
Method EPA 502.2 continued			
tert-Butylbenzene	50-150	30	0.5
Carbon tetrachloride	70-140	30	0.5
Chlorobenzene	37-160	30	0.5
Chloroethane	14-320	30	0.5
Chloroform	51-128	30	0.5
Chloromethane	D-273	30	0.5
2-Chlorotoluene	50-150	30	0.5
4-Chlorotoluene	50-150	30	0.5
DBCP	50-150	30	0.5
Dibromochloromethane	53-149	30	0.5
1,2-Dibromoethane	50-150	30	0.5
Dibromomethane	50-150	30	0.5
1,2-Dichlorobenzene	50-150	30	0.5
1,3-Dichlorobenzene	50-150	30	0.5
1,4-Dichlorobenzene	50-150	30	0.5
Dichlorodifluoromethane	50-150	30	0.5
1,1-Dichloroethane	59-155	30	0.5
1,2-Dichloroethane	49-155	30	0.5
1,1-Dichloroethylene	D-234	30	0.5
cis-1,2-Dichloroethylene	50-150	30	0.5
trans-1,2-Dichloroethylene	54-156	30	0.5
1,2-Dichloropropane	D-210	30	0.5
1,3-Dichloropropane	50-150	30	0.5
2,2-Dichloropropane	50-150	30	0.5
1,1-Dichloropropene	50-150	30	0.5
cis-1,3-Dichloropropene	D-227	30	0.5
trans-1,3-Dichloropropene	17-183	30	0.5
Ethylbenzene	37-162	30	0.5
Hexachlorobutadiene	50-150	30	0.5
Isopropylbenzene	50-150	30	0.5
p-Isopropyltoluene	50-150	30	0.5
Methylene Chloride	D-221	30	0.5
Naphthalene	50-150	30	0.5
n-Propylbenzene	50-150	30	0.5
Styrene	50-150	30	0.5
1,1,1,2-Tetrachloroethane	50-150	30	0.5
1,1,2,2-Tetrachloroethane	46-157	30	0.5
Tetrachloroethylene	64-148	30	0.5
Toluene	47-163	30	0.5
1,2,3-Trichlorobenzene	50-150	30	0.5
1,2,4-Trichlorobenzene	50-150	30	0.5
1,1,1-Trichloroethane	52-150	30	0.5
1,1,2-Trichloroethane	71-157	30	0.5
Trichloroethylene	71-157	30	0.5
Trichlorofluoromethane	17-181	30	0.5

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TABLE 5-1 Quality Assurance Objectives for Drinking Water Methods

	ACCURACY	PRECISION	DLR
CONSTITUENT	% REC-AR	RPD-MAV	ug/L
Method EPA 502.2 continued			
1,2,3-Trichloropropane	50-150	30	0.5
1,1,2-Trichlorotrifluoroeth	50-150	30	0.5
1,2,4-Trimethylbenzene	52-150	30	0.5
1,3,5-Trimethylbenzene	50-150	30	0.5
Vinyl Chloride	D-251	30	0.5
Xylenes m,p	50-150	30	0.5
Xylenes o	50-150	30	0.5
Aylenes	30-130	30	0.3
Method EPA 504			
DBCP	70-130	30	0.02
EDB	70-130	30	0.01
Method EPA 505			
Alachlor	50-150	30	0.2
Aldrin	42-122	30	0.01
Chlordane	45-119	30	0.1
Dieldrin	36-146	30	0.01
Endrin	30-147	30	0.01
Heptachlor	34-111	30	0.01
Heptachlor Epoxide	37-142	30	0.01
Hexachlorobenzene	50-150	30	0.01
Lindane	32-127	30	0.05
Methoxychlor	50-150	30	0.05
Toxaphene	41-126	30	0.5
PCB 1016	50-114	30	0.3
PCB 1221	15-178	30	0.3
PCB 1232	10-215	30	0.3
PCB 1242	39-150	30	0.3
PCB 1248	38-158	30	0.3
PCB 1254	29-131	30	0.3
PCB 1260	8-127	30	0.3
Method EPA 507			
Surrogates			
1,3-Dimethyl-2-nitrobenzene	52 105	NT/A	BT/A
9-Nitroanthracene	53-105	N/A	N/A
Analytes	50-134	N/A	N/A
Alachlor	70.120	20	
Atrazine	70-130 70-130	30	1
Bromocil	70-130	30	1
Butachlor	70-130	30	5
Diazinon	70-130	30	l
Dimethoate	70-130	30	1 2 2
	70-130	30	
Metolachlor Metribusin	70-130	30	1
Metribuzin	70-130	30	0.1

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TABLE 5-1 Quality Assurance Objectives for Drinking Water Methods

CONSTITUENT	ACCURACY % REC-AR	PRECISION RPD-MAV	DLR ug/L
Method EPA 507 continued			
Molinate	70-130	30	2
Prometryne	70-130	30	2 2 1
Propachlor	70-130	30	ī
Simazine	70-130	30	î
Thiobencarb	70-130	30	î
Method EPA 508			
Surrogate			
Hexachlorobenzene	70-130	N/A	N/A
Analytes			
Chlorothalonil	70-130	30	0.2
PCB 1016	50-114	30	0.08
PCB 1221	15-178	30	0.2
PCB 1232	10-215	30	0.2
PCB 1242	39-150	30	0.2
PCB 1248	38-158	30	0.1
PCB 1254	29-131	30	0.1
PCB 1260	8-127	30	0.2
Method EPA 508A			
PCB's as Decachlorobiphenyl	70-130	30	0.2
Method EPA 510			
Bromodichloromethane	70-130	30	0.5
Bromoform	70-130	30	0.5
Chloroform	70-130	30	0.5
Dibromochloromethane	70-130	30	0.5
Method EPA 515.1			
Surrogate	20.150	27/4	N 7/A
2,4-DCAA	30-150	N/A	N/A
Analytes	20.150	20	•
Bentazon	30-150	30	2
Chloramben	30-150	30	1
2,4-D	30-150	30	2
2,4-DB	30-150	30	2
Dalapon	30-150	30	2
Dicamba	30-150	30	2 2 2 5 2
Dichloroprop	30-150	30	
Dinoseb	30-150 30-150	30	1
Pentachlorophenol	30-150	30	0.2
Picloram	30-150 30-150	30	1
2,4,5-T	30-150 30-150	30	1 1
2,4,5-TP (Silvex)	30-150	30	1

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TABLE 5-1 Quality Assurance Objectives for Drinking Water Methods

ACCURACY PRECISION DLR % REC-AR RPD-MAV ug/L Method EPA 524.2 Surrogates 1,2-Dichloroethane-d4 76-114 N/A N/A
Surrogates 1,2-Dichloroethane-d4 76-114 N/A N/A
1,2-Dichloroethane-d4 76-114 N/A N/A
Toluene-d8 88-110 N/A N/A
BFB 86-115 N/A N/A
Analytes
Acetone 50-150 30 0.5
Benzene 37-151 30 0.5
Bromobenzene 50-150 30 0.5
Bromochloromethane 50-150 30 0.5
Bromodichloromethane 35-155 30 0.5
Bromoform 45-169 30 0.5
Bromomethane D-242 30 0.5
2-Butanone (MEK) 50-150 30 0.5
n-Butylbenzene 50-150 30 0.5
sec-Butylbenzene 50-150 30 0.5
tert-Butylbenzene 50-150 30 0.5
Carbon disulfide 50-150 30 0.5
Carbon tetrachloride 70-140 30 0.5
Chlorobenzene 37-160 30 0.5
Chloroethane 14-230 30 0.5
Chloroform 51-138 30 0.5
Chloromethane D-273 30 0.5
2-Chlorotoluene 50-150 30 0.5
4-Chlorotoluene 50-150 30 0.5
Dibromochloromethane 53-149 30 0.5
1,2-Dibromoethane (EDB) 50-150 30 0.5
Dibromomethane 50-150 30 0.5
1,2-Dibromo-3-chloropropane 50-150 30 0.5
1,2-Dichlorobenzene 50-150 30 0.5
1,3-Dichlorobenzene 50-150 30 0.5
1,4-Dichlorobenzene 50-150 30 0.5
Dichlorodifluoromethane 50-150 30 0.5
1,1-Dichloroethane 59-155 30 0.5
1,2-Dichloroethane 49-155 30 0.5
1,1-Dichloroethylene D-234 30 0.5
cis-1,2-Dichloroethylene 50-150 30 0.5
trans-1,2-Dichloroethylene 54-156 30 0.5
1,2-Dichloropropane D-210 30 0.5
1,3-Dichloropropane 50-150 30 0.5
2,2-Dichloropropane 50-150 30 0.5
1,1-Dichloropropene 50-150 30 0.5
cis-1,3-Dichloropropene D-227 30 0.5
trans-1,3-Dichloropropene 17-183 30 0.5
Ethylbenzene 37-162 30 0.5
Hexachlorobutadiene 50-150 30 0.5

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TABLE 5-1 Quality Assurance Objectives for Drinking Water Methods

	ACCURACY	PRECISION	DLR
CONSTITUENT	% REC-AR	RPD-MAV	ug/L
00.1011101111	70 RDC 11R	INI D-MINY	<u>ug/Li</u>
Method EPA 524.2 continued			
2-Hexanone	50-150	30	0.5
Isopropylbenzene	50-150	30	0.5
p-Isopropyltoluene	50-150	30	0.5
Methylene chloride	D-221	30	0.5
4-Methyl-2-pentanone (MIBK)	50-150	30	0.5
Naphthalene	50-150	30	0.5
n-Propylbenzene	50-150	30	0.5
Styrene	50-150	30	0.5
1,1,1,2-Tetrachloroethane	50-150	30	0.5
1,1,2,2-Tetrachloroethane	46-157	30	0.5
Tetrachloroethylene	64-148	30	0.5
Toluene	47-163	30	0.5
1,2,3-Trichlorobenzene	50-150	30	0.5
1,2,4-Trichlorobenzene	50-150	30	0.5
1,1,1-Trichloroethane	52-162	30	0.5
1,1,2-Trichloroethane	52-150	30	0.5
Trichloroethylene	71-157	30	0.5
Trichlorofluoromethane	17-181	30	0.5
1,2,3-Trichloropropane	50-150	30	0.5
1,1,2-Trichlorotrifluoroeth	50-150	30	0.5
1,2,4-Trimethylbenzene	50-150	30	0.5
Vinyl acetate	50-150	30	0.5و
Vinyl chloride	D-251	30	0.5
Xylenes m,p	50-150	30	0.5
Xylenes o	50-150	30	0.5
15 / 17m / 225			
Method EPA 525			
Surrogate	70 1 7 0	37 / 4	37/1
Perylene-d12	50-150	N/A	N/A
Analytes	50 150	20	
Acenaphthylene	50-150	30	1
Anthracene	50-150	30	1
Benzo(a)anthracene	50-150	30	1
Benzo(b)fluoranthene	50-150	30	1
Benzo(k)fluoranthene	50-150	30	1
Benzo(g,h,i)perylene	50-150	30	1
Benzo(a)pyrene	50-150	30	0.1
Butylbenzylphthalate	50-150	30	1
Chrysene	50-150	30	1
Dibenzo(a,h)anthracene	50-150	30	1
Dimethylphthalate	50-150	30	1
Diethylphthalate	50-150	30	1
Di-n-butylphthalate	50-150	30	1
bis(2-Ethylhexyl)adipate	50-150	30	1
bis(2-Ethylhexyl)phthalate	29-137	30	3

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TABLE 5-1 Quality Assurance Objectives for Drinking Water Methods

	ACCURACY	PRECISION	DLR
CONSTITUENT	% REC-AR	RPD-MAV	ug/L
Method EPA 525 continued			
Fluorene	50-150	30	1
Hexachlorobenzene	50-150	30	1
Hexachlorocyclopentadiene	50-150	30	1
Indeno(1,2,3-c,d)pyrene	50-150	30	1
Pentachlorophenol	50-150	30	4
Phenanthrene	50-150	30	1
Pyrene	50-150	30	1
Method EPA 531.1			
Surrogate			
BDMC	70-130	N/A	N/A
Analytes			
Aldicarb	70-130	30	3
Aldicarb Sulfone	70-130	30	3 3 5 5
Aldicarb Sulfoxide	70-130	30	3
Carbofuran	70-130	30	5
Carbaryl	70-130	30	
3-Hydroxycarbofuran	70-130	30	10
Methiocarb	70-130	30	10
Methomyl	70-130	30	5 5 5
1-Napthol	70-130	30	5
Oxymal	70-130	30	5
Propoxur	70-130	30	5
Method EPA 547			
Glyphosate	70-130	20	20
Method EPA 548			
Endothall	70-130	20	40
	70 150	20	40
Method EPA 549			
Diquat	70-130	20	2
Paraquat	70-130	20	1
Method EPA 550.1			
Acenaphthene	70-130	20	3
Acenaphthylene	70-130	20	3 2
Anthracene	70-130	20	0.1
Benzo(a)anthracene	70-130	20	0.1
Benzo(a)pyrene	70-130	20	0.1
Benzo(b)fluoranthene	70-130	20	0.2
Benzo(g,h,i)perylene	70-130	20	0.1
Benzo(k)fluoranthene	70-130	20	0.1
Chrysene	70-130	20	0.1
Dibenzo(a,h)anthracene	70-130	20	0.3

TABLE 5-1 Quality Assurance Objectives for Drinking Water Methods

CONSTITUENT	ACCURACY <u>% REC-AR</u>	PRECISION RPD-MAV	DLR ug/L
Method EPA 550.1 continued			
Fluoranthene	70-130	20	2
Fluorene	70-130	20	2
Indeno(1,2,3-c,d)pyrene	70-130	20	0.1
1-Methylnaphthalene	70-130	20	2
2-Methylnaphthalene	70-130	20	2
Naphthalene	70-130	20	2
Phenanthrene	70-130	20	2
Pyrene	70-130	20	0.1
Method EPA 552			
Bromochloroacetic acid	70-130	20	1
Dibromoacetic acid	70-130	20	1
Dichloroacetic acid	70-130	20	1
2,4-Dichlorophenol	70-130	20	1
Monobromoacetic acid	70-130	20	1
Monochloroacetic acid	70-130	20	1
Trichloroacetic acid	70-130	20	1
2,4,6-Trichlorophenol	70-130	20	1

CONSTITUENT	Method	ACCURACY <u>% REC-AR</u>	PRECISION RPD-MAV	DLR mg/L
Inorganic Chemicals				
Acidity	305.1	N/A	20	1
Alkalinity (as CaCO3)	310.0	N/A	20	1
Bicarbonate	310.1	N/A	20	1
BOD	405.1	80-120	20	2
Bromide	300.0	80-120	20	0.5
Carbon Dioxide	SM4500C	N/A	20	1
Carbonate	310.1	N/A	20	1
COD	410.4	75-125	20	4
Chloride	300.0	80-120	20	1
Chlorine Residual	330.2	N/A	20	0.1
Chlorine Residual	330.5	N/A	20	0.1
Color	110.3	N/A	20	3 units
Electrical Conductivity	120.1	80-120	20	1 umhos
Cyanide, Total	335.2	75-125	20	0.01
Fluoride by electrode	340.2	80-120	20	0.1
Hydroxide	310.0	N/A	20	1
MBAS	425.1	70-130	20	0.05
Nitrogen				
Ammonia	350.1	80-120	20	1
Nitrate	300.0	80-120	20	0.1
Nitrite	300.0	80-120	20	0.1

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TABLE 5-1 Quality Assurance Objectives for Drinking Water Methods

CONSTITUTION	Made a	ACCURACY	PRECISION	DLR
<u>CONSTITUENT</u>	Method	% REC-AR	RPD-MAV	mg/L
Inorganic Chemical EPA Met	hods continued			
Nitrate	353.2	80-120	20	0.1
Nitrite	353.2	80-120	20	0.1
Total Kjeldahl	351.2	80-120	20	1
Odor	140.1	N/A	20	î TON
Oil and Grease	413.1	N/A	20	3
Oxygen, dissolved	360.1	N/A	20	0.5
рН	150.1	N/A	20	N/A
Phenols	420.1	75-125	20	0.1
Phosphorous		, c 120		0.1
Phosphate	300.0	80-120	20	0.1
Total	365.2	75-125	20	0.1
Solids/Residue	500.2	70 120	20	0.1
Filterable (TDS)	160.1	NA	20	40
Non-filterable (TSS)	160.2	NA	20	10
(182)	100.2	IVA.	20	10
Total	160.3	NA	20	40
Volatile	160.4	NA	20	40
Settleable	160.5	NA	20	0.1 ml/L
Sulfate	300.0	80-120	20	1
Sulfide				•
Total	376.2	N/A	20	0.1
Dissolved	376.2	N/A	20	0.1Sulfite
	377.1	N/A	20	0.1
Tannin & Lignin	SM5500B	N/A	20	1
Titration - pH adjust.	N/A	N/A	20	î
Turbidity	180.1	N/A	20	0.2 NTU
·				0.2 1110
G0110777777		ACCURACY	PRECISION	DLR
<u>CONSTITUENT</u>	<u>Method</u>	% REC-AR	RPD-MAV	ug/L
Trace Metals				
Aluminum	200.0	## 10#	20	
	200.9	75-125	20	20
Aluminum Calcium	200.8	75-125	20	20
Chromium	200.7	80-120	20	1 mg/L
Chromium	200.9	75-125	20	5 2
Chromium VI	200.8	75-125	20	
Cobalt	7196	D-120	20	10
	200.7	80-120	20	50
Cobalt	200.8	75-125	20	0.5
Copper	200.7	80-120	20	50
Copper	200.8	75-125	20	2
Iron	200.7	80-120	20	50
Lead	200.9	75-125	20	5
Lead	200.8	75-125	20	0.5
Lithium	SM3500L	80-120	20	10

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TABLE 5-1 Quality Assurance Objectives for Drinking Water Methods

CONSTITUENT	<u>Method</u>	ACCURACY <u>% REC-AR</u>	PRECISION <u>RPD-MAV</u>	DLR ug/L
Trace Metals continued				
Magnesium	200.7	80-120	20	1 mg/L
Manganese	200.7	80-120	20	30
Manganese	200.8	75-125	20	1
Mercury	245.1	75-125	20	0.2
Mercury	245.2	75-125	20	0.2
Molybdenum	200.7	80-120	20	50
Molybdenum	200.8	75-125	20	0.5
Nickel	200.7	80-120	20	50
Nickel	200.8	75-125	20	1
Potassium	200.7	80-120	20	1 mg/L
Selenium	200.9	75-125	20	2 2
Selenium	200.8	75-125	20	2
Silica	200.7	80-120	20	1 mg/L
Silver	200.9	75-125	20	0.5
Silver	200.8	75-125	20	0.5
Sodium	200.7	80-120	20	1 mg/L
Strontium	200.7	80-120	20	50
Thallium	200.9	75-125	20	2
Thallium	200.8	75-125	20	0.5
Tin	200.9	75-125	20	50
Uranium	200.7	80-120	20	100
Vanadium	200.7	80-120	20	20
Zinc	200.7	80-120	20	50
Zinc	200.8	75-125	20	5
		ACCURACY	PRECISION	MDA
CONSTITUENT	<u>Method</u>	% REC-AR	RPD-MAV	pCi/L
Radiochemistry				
Gamma Emitters	901.1	80-120	20	1
Gross Alpha	900.0	60-140	30	1
Gross Beta	900.0	0-140	30	1
Radon	913.0	N/A	20	10
Strontium 90	905.0	60-140	30	1
Total Radium	900.1	75-125	20	1
Radium 226	903.1	75-125	20	1
Radium 228	904.0	75 -12 5	20	ī
Tritium	906.0	75-125	20	300
Uranium	908.0	75-125 75-125	20	1
O' HILLIAM	7000	, J-125		-

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TABLE 5-2 Quality Assurance Objectives for Wastewater / Hazardous Waste Liquid Methods

CONSTITUENT	ACCURACY <u>% REC-AR</u>	PRECISION RPD-MAV	DLR ug/L
M-41 - 1 FD 4 (01/0010			
Method EPA 601/8010			
Surrogates	50 150	B1/4	
BFB	50-150	N/A	N/A
Fluorobenzene	50-150	N/A	N/A
Chlorofluorobenzene	50-150	N/A	N/A
Analytes	10 150	•	
Bromodichloromethane	42-172	20	0.5
Bromoform	13-159	20	0.5
Bromomethane	D-144	20	0.5
Carbon tetrachloride	43-143	20	0.5
Chlorobenzene	38-150	20	0.5
Chloroftane	46-137	20	0.5
Chloroform	49-133	20	0.5
Chloromethane	D-193	20	0.5
Dibromochloromethane	24-191	20	0.5
1,2-Dichlorobenzene	D-208	20	0.5
1,3-Dichlorobenzene	7-187	20	0.5
1,4-Dichlorobenzene	42-143	20	0.5
Dichlorodifluoromethane	50-150	20	0.5
1,1-Dichloroethane	47-132	20	0.5
1,2-Dichloroethane	51-147	20	0.5
1,1-Dichloroethylene	28-167	20	0.5
trans-1,2-Dichloroethylene	38-155	20	0.5
1,2-Dichloropropane	44-156	20	0.5
cis-1,3-Dichloropropene	22-178	20	0.5
trans-1,3-Dichloropropene	22-178	20	0.5
Methylene chloride	25-162	20	0.5
1,1,2,2-Tetrachloroethane	50-150	20	0.5
Tetrachloroethylene	26-162	20	0.5
1,1,1-Trichloroethane	41-138	20	0.5
1,1,2-Trichloroethane	39-139	20	0.5
Trichloroethylene	81-119	20	0.5
Trichlorofluoromethane	21-156	20	0.5
Vinyl chloride	28-163	20	0.5
Method EPA 8015M TPH (purgeable)			
Gas	50-150	40.0	500
Method EPA 8015M TPH (extractable)			
Crude oil	50-150	40.0	500
Diesel fuel	50-150	40.0	500 500
Hydraulic oil	50-150	40.0	500 500
Jet fuel	50-150	40.0	
Stoddard solvent	50-150		500 500
Waste oil	50-150 50-150	40.0	500
·· work VII	3 U-13U	40.0	2000

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TABLE 5-2 Quality Assurance Objectives for Wastewater / Hazardous Waste Liquid Methods

•			•
	ACCURACY	PRECISION	DLR
<u>CONSTITUENT</u>	% REC-AR	RPD-MAV	ug/L
Method EPA 602/8020			
Surrogates	#0 1 #0	~~ .	
BFB	50-150	N/A	N/A
Analytes	50.100	••	
Benzene	78-133	20	0.3
Chlorobenzene	73-132	20	0.3
1,2-Dichlorobenzene	64-143	20	0.3
1,3-Dichlorobenzene	61-150	20	0.3
1,4-Dichlorobenzene	61-151	20	0.3
Ethylbenzene	75-129	20	0.3
Toluene	61-164	20	0.3
Xylene, o	69-137	20	0.3
Xylene, p	71-135	20	0.3
Xylene, m	68-133	20	0.3
N. (1. 1. T.D.). (0.4/00.40 / 1. 1.1.1	.1 . 1 . ()		
Method EPA 604/8040 (analyzed by me		••	4.0
2-Chlorophenol	23-134	30	10
2,4-Dichlorophenol	39-135	30	10
2,4-Dimethylphenol	42-109	30	10
4,6-Dinitro-o-cresol	D-181	30	50
2,4-Dinitrophenol	D-191	30	50
2-Methylphenol	50-150	30	10
4-Methylphenol	50-150	30	10
2-Nitrophenol	29-182	30	10
4-Nitrophenol	29-182	30	50
p-Chloro-m-cresol	22-147	30	20
Pentachlorophenol	14-176	30	50
Phenol	5-112	30	10
2,4,5-Trichlorophenol	37-144	30	10
2,4,6-Trichlorophenol	37-144	30	10
Mathad ED A (00/0000			
Method EPA 608/8080			
Surrogates	26 116	TAT / A	TAT / A
Hexachlorobenzene	26-116	N/A	N/A
Dibutylchlorendate	44-125	N/A	N/A
Analytes	20 122	20	0.3
Aldrin	20-123	30	0.2
Alpha BHC	37-134	30	0.2
Beta BHC	17-147	30	0.2
Delta BHC	19-140	30	0.2
Chlordane	45-119	30	0.2
o,p - DDD	31-141	30	0.2
p,p - DDD	31-141	30	0.2
o,p - DDE	30-145	30	0.2
p,p - DDE	30-145	30	0.2
o,p - DDT	25-160	30	0.2

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TABLE 5-2 Quality Assurance Objectives for Wastewater / Hazardous Waste Liquid Methods

	ACCURACY	PRECISION	DLR
<u>CONSTITUENT</u>	% REC-AR	RPD-MAV	ug/L
M. (1. 1. 177) (00/1000)			
Method EPA 608/8080 continued	5 0.160	20	0.2
p,p - DDT	70-160	30	0.2
Dieldrin	47-126	30	0.2
Endosulfan I	45-153	30	0.2
Endosulfan II	D-202	30	0.2
Endosulfan sulfate	26-144	30	0.2
Endrin	30-147	30	0.2
Endrin aldehyde	50-150	30	0.2
Heptachlor	25-133	30	0.2
Heptachlor epoxide	37-142	30	0.2
Lindane	81-119	30	0.2
Methoxychlor	50-150	30	0.5
Toxaphene	41-126	30	5
PCB 1016	50-114	30	2
PCB 1221	15-178	30	2
PCB 1232	10-215	30	2
PCB 1242	39-150	30	2
PCB 1248	38-158	30	2
PCB 1254	29-131	30	2
PCB 1260	8-127	30	2 2 2 2 2
			-
Method EPA 610/8310			
Acenaphthene	70-130	20	3
Acenaphthylene	70-130	20	3 2
Anthracene	70-130	20	0.1
Benzo(a)anthracene	70-130	20	0.1
Benzo(b)fluoranthene	70-130	20	0.2
Benzo(k)fluoranthene	70-130	20	0.1
Benzo(g,h,i)perylene	70-130	20	0.1
Benzo(a)pyrene	70-130	20	0.2
Chrysene	70-130	20	0.1
Dibenzo(a,h)anthracene	70-130	20	0.3
Fluoranthene	70-130	20	2
Fluorene	70-130	20	2
Indeno(1,2,3-c,d)pyrene	70-130	20	0.1
Naphthalene	70-130	20	
Phenanthrene	70-130	20	2 2
Pyrene	70-130 70-130	20 20	0.1
Tyrene	/0-130	20	0.1
Method EPA 614/8140			
Surrogates			
1,3-Dimethyl-2-nitrobenzene	50 150	NI/A	NT/A
9-Nitroanthracene	50-150	N/A	N/A
Analytes	50-150	N/A	N/A
	50.150	20	•
Azinphos methyl Bolstar	50-150	30	2
DUISTAL	50-150	30	2

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TABLE 5-2 Quality Assurance Objectives for Wastewater / Hazardous Waste Liquid Methods

	ACCUDACY	DDECICION	DID
CONSTITUENT	ACCURACY % REC-AR	PRECISION <u>RPD-MAV</u>	DLR
CONSTITUENT	70 KEC-AK	KI D-MAY	ug/L
Method EPA 614/8140 continued			
Chlorpyrifos	50-150	30	2
Coumaphos	50-150	30	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2
Demeton-o,s	50-150	30	2
Diazinon	50-150	30	2
Dichlorvos	50-150	30	2
Disulfoton	50-150	30	2
Ethion	50-150	30	2
Ethoprop	50-150	30	2
Fensulfoton	50-150	30`	2
Fenthion	50-150	30	2
Malathion	50-150	30	2
Merphos	50-150	30	2
Mevinphos	50-150	30	2
Naled	50-150	30	2
Parathion, ethyl	50-150	30	2
Parathion, methyl	50-150	30	2
Phorate	50-150	30	2
Ronnel	50-150	30	2
Stirophos	50-150	30	2
Tokuthion	50-150	30	2
Trichlornate	50-150	30	2
Method EPA 615/8150			
Surrogate			
2,4-DCAA	30-150	N/A	N/A
Analytes			
Bentazon	30-150	30	20
Chloramben	30-150	30	10
2,4-D	30-150	30	100
Dalapon	30-150	30	10
2,4-DB	30-150	30	100
Dicamba	30-150	30	10
Dichlorprop	30-150	30	20
Dinoseb	30-150	30	10
Pentachlorophenol	30-150	30	10
Picloram	30-150	30	10
2,4,5-T	30-150	30	10
2,4,5-TP (Silvex)	30-150	30	10
Method EPA 624/8240			
Surrogates			
1,2-Dichloroethane-d4	76-114	30	N/A
Toluene-d8	88-110	30	N/A
BFB	86-115	30	N/A

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TABLE 5-2 Quality Assurance Objectives for Wastewater / Hazardous Waste Liquid Methods

CONSTITUENT	ACCURACY <u>% REC-AR</u>	PRECISION <u>RPD-MAV</u>	DLR ug/L
Method EPA 624/8240 continued			
Analytes			
Acetone	50-150	30	10
Acrolein	50-150	30	100
Acrylonitrile	50-150	30	100
Benzene	37-151	30	0.5
Bromodichloromethane	35-155	30	1
Bromoform	45-169	30	ī
Bromomethane	D-242	30	ī
Carbon disulfide	50-150	30	5
Carbon tetrachloride	70-140	30	0.5
Chlorobenzene	37-160	30	0.5
Chloroethane	14-230	30	1
2-Chloroethylvinyl ether	50-150	30	10
Chloroform	51-138	30	0.5
Chloromethane	D-273	30	1
Dibromochloromethane	53-149	30	1
1,2-Dichlorobenzene	50-150	30	1
1,3-Dichlorobenzene	50-150	30	1
1,4-Dichlorobenzene	50-150	30	1
Dichlorodifuoromethane	50-150	30	0.5
1,1-Dichloroethane	59-155	30	1
1,2-Dichloroethane	49-155	30	1
1,1-Dichloroethylene	D-234	30	1
trans-1,2-Dichloroethylene	54-156	30	1
1,2-Dichloropropane	D-210	30	1
cis-1,3-Dichloropropene	D-227	30	2
trans-1,3-Dichloropropene	17-183	30	1
Ethanol	50-150	30	5000.
Ethylbenzene	37-162	30	0.5
2-Hexanone	50-150	30	5
Methylene chloride	D-221	30	0.5
2-Butanone (MEK)	50-150	30	10
4-Methyl-2-pentanone (MIBK) Styrene	50-150	30	5
1,1,2,2-Tetrachloroethane	50-150	30	1
Tetrachloroethylene	46-157	30	1
Toluene	64-148	30	1
1,1,1-Trichloroethane	47-163 52-163	30	0.5
1,1,2-Trichloroethane	52-162 53-150	30	0.5
Trichloroethylene	52-150 71 157	30	0.5
Trichlorofluoromethane	71-157	30	1
Vinyl acetate	17-181	30	1.5
Vinyl chloride	50-150 D 251	30	100
Xylenes	D-251	30	0.5
-ry relies	50-150	30	1

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TABLE 5-2 Quality Assurance Objectives for Wastewater / Hazardous Waste Liquid Methods

-			_
	ACCURACY	PRECISION	DLR
<u>CONSTITUENT</u>	<u>% REC-AR</u>	<u>RPD-MAV</u>	<u>ug/L</u>
Method EPA 625/8270			
Surrogates		••	
2-Fluorobiphenyl	43-106	30	N/A
Nitrobenzene-d5	39-98	30	N/A
p-Terphenyl-d14	62-114	30	N/A
2-Fluorophenol	21-100	30	N/A
Phenol-d6	10-94	30	N/A
2,4,6-Tribromophenol	11-100	30	N/A
Analytes			
Acenaphthene	36-130	30	10
Acenaphthylene	33-145	30	10
Aniline	50-150	30	50
Anthracene	27-133	30	10
Benzo(a)anthracene	33-143	30	10
Benzo(a)pyrene	17-163	30	10
Benzo(b)fluoranthene	24-159	30	10
Benzo(k)fluoranthene	11-162	30	10
Benzo(g,h,i)perylene	D-219	30	10
Benzylalcohol	50-150	30	20
bis(2-Chloroethoxy)methane	33-184	30	10
bis(2-Chloroethyl)ether	12-158	30	10
bis(2-Chloroisopropyl)ether	36-166	30	10
bis(2-Ethylhexly)phthalate	29-137	30	10
4-Bromophenylphenylether	65-114	30	10
Butylbenzylphthalate	D-152	30	10
Chloroaniline	50-150	30	10
Chloronaphthalene	60-180	30	10
Chlorophenylphenylether	25-158	30	10
Chrysene	17-168	30	10
Dibenzo(a,h)anthracene	D-227	30	10
Dibenzofuran	50-150	30	10
1,2-Dichlorobenzene	32-129	30	10
1,3-Dichlorobenzene	D-172	30	10
1,4-Dichlorobenzene	33-114	30	10
3,3'-Dichlorbenzidine	8-213	30	20
Diethylphthalate	D-114	30	10
Dimethylphthalate	D-112	30	10
Di-n-butylphthalate	1-118	30	10
2,4-Dinitrotoluene	37-111	30	10
2,6-Dinitrotoluene	50-158	30	10
Di-n-octylphthalate	4-146	30	10
Fluoranthene	26-137	30	10
Fluorene	59-121	30	10
Hexachlorobenzene	50-150	30	10
Hexachlorobutadiene	24-116	30	10
Hexachlorocyclopentadiene	50-150	30	10
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TABLE 5-2 Quality Assurance Objectives for Wastewater / Hazardous Waste Liquid Methods

	ACCURACY	PRECISION	DLR
CONSTITUENT	% REC-AR	RPD-MAV	ug/L
Method EPA 625/8270 continued			
Hexachloroethane	40-113	20	10
		30 30	10
Indeno(1,2,3-c,d)pyrene Isophorone	50-150 21-196	30 30	10
2-Methylnaphthalene	50-150	30 30	10
Naphthalene	21-133	30 30	10
Nitrobenzene	35-180	30 30	10
N-Nitrosodimethylamine	50-150	30 30	10 10
N-Nitrosodi-N-propylamine	D-112	30	10
N-Nitrosodiphenylamine	50-150	30	10
2-Nitroanaline	50-150	30	50
3-Nitroanaline	50-150	30	50 50
4-Nitroanaline	50-150	30	50 50
Phenanthrene	54-120	30	10
Pyrene	73-119	30	10
1,2,4-Trichlorbenzene	35-122	30	10
2-Chlorophenol	27-143	30	10
2,4-Dichlorophenol	39-135	30	10
2,4-Dimethylphenol	42-109	30	10
4,6-Dinitro-o-cresol	D-181	30	50
2,4-Dinitrophenol	D-101 D-191	30	50 50
2-Methylphenol	50-150	30	10
4-Methylphenol	50-150	30	10
2-Nitrophenol	29-182	30	10
4-Nitrophenol	D-53	50	50
p-Chloro-m-cresol	22-147	30	20
Pentachlorophenol	D-112	30	50
Phenol	D-82	30	10
2,4,5-Trichlorophenol	50-150	30	10
2,4,6-Trichlorophenol	37-144	30	10
•			10
Method EPA 632			
Barban	30-150	30	0.5
Carbaryl	30-150	30	0.2
Carbofuran	30-150	30	30
Chlorpropham	30-150	30	0.3
Diuron	30-150	30	0.1
Fluometuron	30-150	30	100
Linoron	30-150	30	0.1
Methiocarb	30-150	30	0.2
Methomyl	30-150	30	100
Monuron	30-150	30	0.1
Neburon	30-150	30	0.1
Oxamyl	30-150	30	100
Propham	30-150	30	0.7
Propoxur	30-150	30	1

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TABLE 5-2 Quality Assurance Objectives for Wastewater / Hazardous Waste Liquid Methods

CONSTITUENT			URACY EC-AR		ECISION D-MAV	DL:	
Method EPA 415.1/9060 TOC		80-12	0	20		0.5	mg/L
Method EPA 9020 TOX		80-12	0	20		0.5	
Method EPA 418.1 TRPH-By IR		50-15	50	20		0.5	mg/L
CONSTITUENT	Method		ACCURACY % REC-AR		PRECISION RPD-MAV		DLR mg/L
Inorganic Chemicals Acidity Alkalinity (as CaCO3) Bicarbonate BOD Bromide Carbon Dioxide Carbonate COD Chloride Chlorine Residual Chlorine Residual Color Electrical Conductivity Cyanide, Total Fluoride by electrode Hydroxide MBAS Nitrogen Ammonia Nitrate Nitrite Nitrite Total Kjeldahl Odor Oil and Grease Oxygen, dissolved	305.1 310.0 310.1 405.1 300.0 SM45000 310.1 410.4 300.0 330.2 330.5 110.3 120.1 335.2 340.2 310.0 425.1 350.1 300.0 300.0 353.2 353.2 353.2 351.2 140.1 413.1 360.1	С	N/A N/A N/A 80-120 80-120 N/A N/A 75-125 80-120 N/A N/A N/A N/A 80-120 75-125 80-120 N/A 70-130 80-120 80-120 80-120 80-120 80-120 80-120 80-120 N/A N/A N/A N/A		20 20 20 20 20 20 20 20 20 20		1 1 1 2 0.5 1 1 4 1 0.1 0.1 3 units 1 umhos 0.01 0.1 1 0.1 0.1 0.1 0.1 0.1 0.1
pH Phenols Phosphorous phosphate Total	150.1 420.1 300.0 365.2		N/A 75-125 80-120 75-125		20 20 20 20 20		N/A 0.1 0.1 0.1

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TABLE 5-2 Quality Assurance Objectives for Wastewater / Hazardous Waste Liquid Methods

CONSTITUENT	Method	ACCURACY % REC-AR	PRECISION RPD-MAV	DLR mg/L
Inorganic Chemicals continue	d			
Solids/Residue				
Filterable (TDS)	160.1	NA	20	40
Non-filterable (TSS)	160.2	NA	20	10
Total	160.3	NA	20	40
Volatile	160.4	NA	20	40
Settleable	160.5	NA	20	0.1 ml/L
Sulfate	300.0	80-120	20	1
Sulfide				_
Total	376.2	N/A	20	0.1
Dissolved	376.2	N/A	20	0.1
Sulfite	377.1	N/A	20	0.1
Tannin & Lignin	SM5500B	N/A	20	1
Titration - pH adjust.	N/A	N/A	20	ĩ
Turbidity	180.1	N/A	20	0.2 NTU
,	20012			0.2 1110
Trace Metals				
Aluminum	200.9	75-125	20	0.02
Aluminum	200.8	75-125	20	0.1
Antimony	200.9	75-125	20	0.02
Antimony	200.8	75-125	20	0.005
Arsenic	200.9	75-125	20	0.005
Arsenic	200.8	75-125	20	0.003
Barium	200.7	80-120	20	0.02
Barium	200.8	75-125	20	0.02
Beryllium	200.7	80-120	20	0.003
Beryllium	200.8	75-125	20	0.01
Boron	200.7	80-120	20	0.003
Boron	200.8	75-125	20	0.1
Cadmium	200.9	75-125 75-125	20	0.005
Cadmium	200.8	75-125 75-125	20	0.005
Calcium	200.7	80-120	20	1
Chromium	200.9	75-125	20	0.01
Chromium	200.8	75-125 75-125	20	0.01
Chromium VI	7196	D-120	20	0.02
Cobalt	200.7	80-120	20	0.05
Cobalt	200.8	75-125	20	0.005
Copper	200.7	80-120	20	0.003
Copper	200.8	75-125	20	
Iron	200.7	80-120	20	0.02
Gold	231.1			0.05
Lead		75-125 75-125	20	0.05
Lead	200.9	75-125 75-125	20	0.01
Organic Lead	200.8	75-125 50 150	20	0.005
Lithium	LUFT	50-150	20	0.05
Magnesium	SM3500L	80-120	20	0.05
Magnesium	200.7	80-120	20	1

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TABLE 5-2 Quality Assurance Objectives for Wastewater / Hazardous Waste Liquid Methods

		ACCURACY	PRECISION	DLR
<u>CONSTITUENT</u>	<u>Method</u>	% REC-AR	RPD-MAV	mg/L
Trace Metals continued		00.100		
Manganese	200.7	80-120	20	0.03
Manganese	200.8	75-125	20	0.01
Mercury	245.1	75-125	20	0.0002
Mercury	245.2	75-125	20	0.0002
Molybdenum	200.7	80-120	20	0.05
Molybdenum	200.8	<i>75-125</i>	20	0.005
Nickel	200.7	80-120	20	0.05
Nickel	200.8	75-125	20	0.01
Potassium	200.7	80-120	20	1
Selenium	200.9	75-125	20	0.005
Selenium	200.8	75-125	20	0.02
Silica	200.7	80-120	20	1
Silver	200.9	75-125	20	0.01
Silver	200.8	75-125	20	0.005
Sodium	200.7	80-120	20	1
Strontium	200.7	80-120	20	0.05
Thallium	200.9	75-125	20	0.02
Thallium	200.8	75-125	20	0.005
Tin	200.9	75-125	20	0.05
Titanium	200.7	80-120	20	0.1
Uranium	200.7	80-120	20	0.1
Vanadium	200.7	80-120	20	0.02
Zinc	200.7	80-120	20	0.05
Zinc	200.8	75-125	20	0.05
Zinc	200.0	75-125	20	0.05
		ACCURACY	PRECISION	MDA
CONSTITUENT	Method	% REC-AR	RPD-MAV	pCi/L
				<u></u>
Radiochemistry				
Gamma Emitters	901.1	80-120	20	1
Gross Alpha	900.0	60-140	30	1
Gross Beta	900.0	60-140	30	1
Radon	913.0	N/A	20	10
Strontium 90	905.0	60-140	30	1
Total Radium	900.1	75-125	20	1
Radium 226	903.1	75-125	20	1
Radium 228	904.0	75-125	20	1
Tritium	906.0	75-125	20	300
Uranium	908.0	75-125	20	1

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TABLE 5-3 Quality Assurance Objectives for Solid Waste / Hazardous Waste Methods

CONSTITUENT	ACCURACY % REC-AR	PRECISION RPD-MAV	DLR mg/Kg
CONSTITUENT	70 REC-AR	KI D-MAY	mg/Kg
Method EPA 8010			
Surrogates			
BFB	50-150	N/A	N/A
Fluorobenzene	50-150	N/A	N/A
Chlorofluorobenzene	50-150	N/A	N/A
Analytes			
Azobenzene	50-150	30	5
Benzidine	50-150	30	5
Benzoic acid	50-150	30	5
Benzylchloride	50-150	20	5 5 5 5 5 5 5
bis(2-Chloroisopropyl)ether	50-150	20	5
Bromobenzene	50-150	20	5
Bromochloromethane	50-150	20	5
Bromodichloromethane	42-172	20	5
Bromoform	13-159	20	5
Bromomethane	D-144	20	5
Carbon tetrachloride	43-143	20	5
Chlorobenzene	38-150	20	5 5 5 5 5
Chloroethane	46-137	20	5
Chloroform	49-133	20	5
1-Chlorohexane	50-150	20	5
Chloromethane	D-193	20	5
2-Chlorotoluene	50-150	20	5
Chlorotoluene	50-150	20	5
DBCP	50-150	20	5
Dibromochloromethane	24-191	20	5
1,2-Dibromoethane	50-150	20	5
Dibromomethane	50-150	20	5
1,2-Dichlorobenzene	D-208	20	5 5 5 5 5
1,3-Dichlorobenzene	7-187	20	5
1,4-Dichlorobenzene	42-143	20	5
1,1-Dichloroethane	47-132	20	5
1,2-Dichloroethane	51-147	20	5
1,1-Dichloroethylene	28-167	20	5
cis-1,2-Dichloroethylene	50-150	20	5
trans-1,2-Dichloroethylene	38-155	20	5
1,2-Dichloropropane	44-156	20	5
1,3-Dichloropropane	50-150	20	5
2,2-Dichloropropane	50-150	20	5
1,1-Dichloropropene	50-150	20	5
cis-1,3-Dichloropropene	22-178	20	5
trans-1,3-Dichloropropene	22-178	20	555555555555
Hexachlorobutadiene	50-150	20	5
Methylene chloride	25-162	20	5
1,1,1,2-Tetrachloroethane	50-150	20	5
1,1,2,2-Tetrachloroethane	50-150	20	5

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TABLE 5-3 Quality Assurance Objectives for Solid Waste / Hazardous Waste Methods

CONSTITUENT	ACCURACY % REC-AR	PRECISION RPD-MAV	DLR mg/Kg
Method EPA 8010 continued			
Tetrachloroethylene	26-162	20	5
1,2,3-Trichlorobenzene	50-150	20	5 5 5 5 5 5 5
1,2,4-Trichlorobenzene	50-150	20	5
1,1,1-Trichloroethane	41-138	20	5
1,1,2-Trichloroethane	39-136	20	5
Trichloroethylene	35-146	20	5
Trichlorofluoromethane	21-156	20	5
Trichloropropane	50-150	20	5
Vinyl chloride	28-163	20	5
Method EPA 8011			
EDB	50-150	30	0.0005
DBCP	50-150	30	0.0005
Method EPA 8015			
Acrylamide	50-150	40	0.05
Diethyl ether	50-150	40	0.05
Ethanol	50-150	40	0.05
Methyl ethyl ketone	50-150	40	0.05
Methyl isobutyl ketone	50-150	40	0.05
Paraldehyde	50-150	40	0.05
Method EPA 8015M TPH (purgeable)			
Gas	50-150	40	5
Method EPA 8015M TPH (extractable)			
Crude oil	50-150	40	10
Diesel fuel	50-150	40	10
Hydraulic oil	50-150	40	10
Jet fuel	50-150	40	10
Stoddard solvent	50-150	40	10
Waste oil	50-150	40	50
Method EPA 8020			
Benzene	39-159	20	0.003
Chlorobenzene	55-135	20	0.003
Ethylbenzene	32-160	20	0.003
1,2-Dichlorobenzene	37-154	20	0.003
1,3-Dichlorobenzene	50-141	20	0.003
1,4-Dichlorobenzene	42-143	20	0.003
Toluene	46-148	20	0.003
Xylene, o	50-150	20	0.003
Xylene, p	50-150	20	0.003
Xylene, m	50-150	20	0.003

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TABLE 5-3 Quality Assurance Objectives for Solid Waste / Hazardous Waste Methods

CONSTITUENT	ACCURACY <u>% REC-AR</u>	PRECISION RPD-MAV	DLR mg/Kg			
Method FPA 80/0 (analyzed by metho	.d 9270)					
Method EPA 8040 (analyzed by method 8270) 2-s-Butyl-4-6-dinitrophenol 50-150 30 0.1						
2-s-Butyl-4-6-dinitrophenol 2-Chlorophenol	23-134	30 30	0.1			
4-Chloro-3-methylphenol	23-134 22-147		0.1			
2,4-Dichlorophenol	39-135	30	0.2			
		30	0.1			
2,4-Dimethylphenol	42-109 D 101	30	0.1			
2,4-Dinitrophenol	D-191	30	0.5			
2-Methyl-4,6-Dinitrophenol	D-181	30	0.5			
2-Methylphenol	50-150	30	0.1			
4-Methylphenol	50-150	30	0.1			
2-Nitrophenol	29-182	30	0.1			
4-Nitrophenol	29-182	30	0.5			
Pentachlorophenol	14-176	30	0.5			
Phenol	5-112	30	0.1			
2,3,4,6-Tetrachlorophenol	50-150	30	0.1			
2,3,5,6-Tetrachlorophenol	50-150	30	0.1			
2,3,4-Trichlorophenol	37-144	30	0.1			
2,3,5-Trichlorophenol	37-144	30	0.1			
2,3,6-Trichlorophenol	37-144	30	0.1			
2,4,5-Trichlorophenol	37-144	30	0.1			
2,4,6-Trichlorophenol	37-144	30	0.1			
Method EPA 8080						
Surrogates						
Hexachlorobenzene	50-150	30	N/A			
Dibutylchlorendate	20-150	30	N/A			
Analytes	20 130	30	IVA			
Aldrin	34-132	43	0.05			
Alpha BHC	37-134	30	0.05			
Beta BHC	17-147	30	0.05			
Delta BHC	19-140	30	0.05			
Chlordane	45-119	30	0.05			
o,p - DDD	31-141	30	0.05			
p,p - DDD	31-141	30	0.05			
o,p - DDE	30-145	30				
p,p - DDE	30-145	30	0.05			
o,p - DDT	23-134	50	0.05			
p,p - DDT	23-134 23-134		0.05			
Dieldrin		50	0.05			
Endosulfan I	31-134	38	0.05			
Endosulfan II	45-153 D 202	30	0.05			
Endosulfan 11 Endosulfan sulfate	D-202	30	0.05			
Endosulian sulfate Endrin	26-144	30	0.05			
	42-139	45	0.05			
Endrin aldehyde	50-150	30	0.05			
Heptachlor	35-130	30	0.05			
Heptachlor epoxide	37-142	30	0.05			

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TABLE 5-3 Quality Assurance Objectives for Solid Waste / Hazardous Waste Methods

	ACCURACY	PRECISION	DLR
CONSTITUENT	% REC-AR	RPD-MAV	mg/Kg
Method EPA 8080 continued			
Lindane	46-127	50	0.05
Methoxychlor	50-150	30	0.1
Toxaphene	41-126	30	0.1
PCB 1016	50-114	30	0.5
PCB 1221	15-178	30	0.5
PCB 1232	10-215	30	0.5
PCB 1242	39-150	30	0.5
PCB 1248	38-158	30	0.5
PCB 1254	29-131	30	0.5
PCB 1260	8-127	30	0.5
Method EPA 8140			
Surrogates			
1,3-Dimethyl-2-nitrobenzene	50-150	30	N/A
9-Nitroanthracene	50-150	30	N/A
Analytes			
Azinphos methyl	50-150	30	0.02
Bolstar	50-150	30	0.02
Chlorpyrifos	50-150	30	0.02
Coumaphos	50-150	30	0.02
Demeton-o,s	50-150	30	0.02
Diazinon	50-150	30	0.02
Dichlorvos	50-150	30	0.02
Disulfoton	50-150	30	0.02
Ethoprop	50-150	30	0.02
Fensulfoton	50-150	30	0.02
Fenthion	50-150	30	0.02
Merphos	50-150	30	0.02
Mevinphos	50-150	30	0.02
Naled	50-150	30	0.02
Parathion methyl	50-150	30	0.02
Phorate	50-150	30	0.02
Ronnel	50-150	30	0.02
Stirophos	50-150	30	0.02
Tokuthion	50-150	30	0.02
Trichlornate	50-150	30	0.02
Method EPA 8150			
Surrogate			
2,4-DČAA	30-150	N/A	N/A
Analytes			
Bentazon	30-150	30	0.2
Chloramben	30-150	30	0.1
2,4-D	30-150	30	1
Dalapon	30-150	30	0.1
-			

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TABLE 5-3 Quality Assurance Objectives for Solid Waste / Hazardous Waste Methods

CONSTITUENT	ACCURACY <u>% REC-AR</u>	PRECISION <u>RPD-MAV</u>	DLR mg/Kg
Method EPA 8150 continued			
2,4-DB	30-150	30	1
Dicamba	30-150	30	1
Dichlorprop	30-150		0.1
Dinoseb	30-150 30-150	30	0.2
Pentachlorophenol		30	0.1
Picloram	30-150	30	0.1
2,4,5-T	30-150	30	0.1
2,4,5-TP (Silvex)	30-150 30-150	30 30	0.1 0.2
Method EPA 8240			
Surrogates			
1,2-Dichloroethane-d4	61-164	N/A	NI/A
Toluene-d8	81-117	N/A N/A	N/A
BFB	67-124		N/A
Analytes	0/-124	N/A	N/A
Acetone	E0 150	20	0.01
Acrolein	50-150	30	0.01
Acrylonitrile	50-150	30	0.1
Benzene	50-150	30	0.1
Bromodichloromethane	66-142	21	0.005
Bromoform	35-155 45-160	30	0.005
Bromomethane	45-169 D. 242	30	0.005
Carbon disulfide	D-242	30	0.01
	50-150	30	0.005
Carbon tetrachloride	70-140	30	0.005
Chlorosthana	60-133	21	0.005
Chloroethane	14-230	30	0.01
2-Chloroethylvinyl ether	50-150	30	0.01
Chloroform Chloromethane	51-138	30	0.005
	D-273	30	0.01
Dibromochloromethane	53-149	30	0.005
1,2-Dichlorobenzene	50-150	30	0.005
1,3-Dichlorobenzene	50-150	30	0.005
1,4-Dichlorobenzene	50-150	30	0.005
Dichlorodifluoromethane	50-150	30	0.0005
1,1-Dichloroethane	59-172	21	0.005
1,2-Dichloroethane	49-155	30	0.005
1,1-Dichloroethylene	D-234	30	0.005
trans-1,2-Dichloroethylene	54-156	30	0.005
1,2-Dichloropropane	D-210	30	0.005
cis-1,3-Dichloropropene	D-227	30	0.005
trans-1,3-Dichloropropene	17-183	30	0.005
Ethanol	50-150	30	10
Ethylbenzene	37-162	30	0.005
2-Hexanone	50-150	30	0.005
Methylene chloride	D-221	30	0.005

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TABLE 5-3 Quality Assurance Objectives for Solid Waste / Hazardous Waste Methods

CONSTITUENT	ACCURACY % REC-AR	PRECISION RPD-MAV	DLR mg/Kg
Method EPA 8240 continued			
2-Butanone (MEK)	50-150	30	0.01
4-Methyl-2-pentanone (MIBK)	50-150	30	0.005
Styrene	50-150	30	0.005
1,1,2,2-Tetrachloroethane	46-157	30	0.005
Tetrachloroethylene	64-148	30	0.005
Toluene	59-139	21	0.005
1,1,1-Trichloroethane	52-162	30	0.005
1,1,2-Trichloroethane	52-150	30	0.005
Trichlorethylene	62-137	24	0.005
Trichlorofluoromethane	17-181	30	0.005
Vinyl acetate	50-150	30	0.1
Vinyl chloride	D-251	30	0.01
Xylenes	50-150	30	0.005
Method EPA 8270 Surrogates			
2-Fluorobiphenyl	30-115	30	N/A
Nitrobenzene-d5	23-120	30	N/A
p-Terphenyl-d14	18-137	30	N/A
2-Fluorophenol	25-121	30	N/A
Phenol-d6	24-113	30	N/A
2,4,6-Tribromophenol	19-122	30	N/A
Analytes			
Acenaphthene	31-137	19	1
Acenaphthylene	33-145	30	1
Aniline	50-150	30	5
Anthracene	27-133	30	1
Benzo(a)anthracene	33-143	30	1
Benzo(a)pyrene	17-163	30	1
Benzo(b)fluoranthene	24-159	30	1
Benzo(k)fluoranthene	11-162	30	1
Benzo(g,h,i)perylene	D-219	30	1
Benzylalcohol	50-150	30	2
bis(2-Chloroethoxy)methane	33-184	30	1
bis(2-Chloroethyl)ether	12-158	30	1
bis(2-Chloroisopropyl)ether	36-166	30	1
bis(2-Ethylhexyl)phthalate	29-137	30	1
4-Bromophenylphenylether	65-114	30	1 .
Butylbenzylphthalate	D-152	30	1
Chloroaniline	50-150	30	2
Chloronaphthalene	60-180	30	1
Chlorophenylphenylether	25-158	30	1
Chrysene	17-168	30	1
Dibenzo(a,h)anthracene	D-227	30	1
Dibenzofuran	50-150	30	1

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TABLE 5-3 Quality Assurance Objectives for Solid Waste / Hazardous Waste Methods

CONSTITUENT	ACCURACY % REC-AR	PRECISION RPD-MAV	DLR mg/Kg
Method EPA 8270 continued			
1,2-Dichlorobenzene	32-129	20	1
		30	1
1,3-Dichlorobenzene	D-172	30	1
1,4-Dichlorobenzene	28-104	27	1
3,3'-Dichlorobenzidine	8-213	30	2
Diethylphthalate	D-114	30	1
Dimethylphthalate	D-112	30	1
Di-n-butylphthalate	1-118	30	1
2,4-Dinitrotoluene	28-100	47	1
2,6-Dinitrotoluene	50-158	30	1
Di-n-octylphthalate	4-146	30	1
Fluoranthene	26-137	30	1
Fluorene	59-121	30	1
Hexachlorobenzene	50-150	30	1
Hexachlorobutadiene	24-116	30	1
Hexachlorocyclopentadiene	50-150	30	2
Hexachloroethane	40-113	30	1
Indeno(1,2,3-c,d)pyrene	50-150	30	1
Isophorone	21-196	30	1
2-Methylnaphthalene	50-150	30	1
Naphthalene	21-133	30	1
Nitrobenzene	35-180	30	1
N-Nitrosodimethylamine	50-150	30	1
N-Nitrosodi-N-propylamine	41-126	38	1
N-Nitrosodiphenylamine	50-150	30	1
2-Nitroanaline	50-150	30	5
3-Nitroanaline	50-150	30	5
4-Nitroanaline	50-150	30	5
Phenanthrene	54-120	30	1
Pyrene	35-142	36	1
1,2,4-Trichlorbenzene	38-107	23	1
2-Chlorophenol	25-102	50	1
2,4-Dichlorophenol	39-135	30	1
2,4-Dimethylphenol	42-109	30	1
4,6-Dinitro-o-cresol	D-181	30	5
2,4-Dinitrophenol	D-191	30	5 5
2-Methylphenol	50-150	30	1
4-Methylphenol	50-150	30	1
2-Nitrophenol	29-182	30	î
4-Nitrophenol	11-114	50	5
p-Chloro-m-cresol	26-103	33	2
Pentachlorophenol	17-109	47	5
Phenol	26-90	35	1
2,4,5-Trichlorophenol	50-150	30	1
2,4,6-Trichlorophenol	37-144	30	1
Azobenzene	50-150	30	5
	20 130	50	5

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TABLE 5-3 Quality Assurance Objectives for Solid Waste / Hazardous Waste Methods

CONSTITUENT		ACCURACY % REC-AR	PRECISION RPD-MAV	DLR mg/Kg
Method EPA 8270 continued Benzidine Benzoic Acid		50-150 50-150	30 30	5 5
Method EPA 8310 Acenaphthene Acenaphthylene Anthracene Benzo(a)anthracene Benzo(b)fluoranthene		70-130 70-130 70-130 70-130 70-130	20 20 20 20 20	3 2 0.1 0.1 0.2
Benzo(k)fluoranthene Benzo(g,h,i)perylene Benzo(a)pyrene Chrysene Dibenzo(a,h)anthracene Fluoranthene		70-130 70-130 70-130 70-130 70-130 70-130	20 20 20 20 20 20 20	0.1 0.1 0.2 0.1 0.3 2
Fluorene Indeno(1,2,3-c,d)pyrene Naphthalene Phenanthrene Pyrene		70-130 70-130 70-130 70-130 70-130	20 20 20 20 20 20	2 0.1 2 2 0.1
Method EPA 9060 TOC		80-120	20	50
Method EPA 9020 TOX		80-120	20	1 .
EPA Method 418.1M TRPH-By IR		50-150	20	10
CONSTITUENT	Method	ACCURACY <u>% REC-AR</u>	PRECISION RPD-MAV	DLR mg/L
Inorganic Chemicals Chloride Electrical Conductivity Cyanide, Total Fluoride	9056 120.1 335.2 335.2	70-130 80-120 65-135 65-135	30 20 30 30	10 1 umhos 1 50

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TABLE 5-3 Quality Assurance Objectives for Solid Waste / Hazardous Waste Methods

CONSTITUENT	Method	ACCURACY <u>% REC-AR</u>	PRECISION RPD-MAV	DLR mg/L
Inorganic Chemicals				
Nitrogen				
Ammonia-N	350.1	70-130	30	4
Nitrate	9056	70-130	30	4
Nitrite	9056	70-130	30	3
Organic	Calc.	N/A	30	100
Kjeldahl	351.1	65-135	30	100
Total	Calc.	N/A	30	100
Moisture	ASA/UL	N/A	20	N/A
Oil and Grease, Soxhlet	413.1	N/A	30	300
рH	150.1	N/A	20	N/A
Phenols	420.1	65-135	30	5
Phosphorous				C
Phosphate	9056	70-130	30	3
Total (see Trace Metals)	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	, , , ,		2
Sulfate	9056	70-130	30	10
Sulfide	376.2	N/A	30	5
	-			J
Hazardous Waste Character	ization			
Corrosivity (pH)	9045	20.0	N/A	N/A
Ignitability "	1020	N/A	N/A	N/A
Reactivity	SW-846	N/A	N/A	N/A
Generation	SW-846	N/A	N/A	N/A
Trace Metals				
Aluminum	6010	70-130	30	50
Aluminum	6020	70-130	30	20
Antimony	7041	65-135	30	3
Antimony	6020	65-135	30	1
Arsenic	7060	65-135	30	1
Arsenic	6020	65-135	30	4
Barium	6010	70-130	30	1
Barium	6020	70-130	30	1
Beryllium	6010	70-130	30	0.5
Beryllium	6020	70-130	30	1
Boron	6010	70-130	30	
Boron	6020	70-130	30	5
Cadmium	6010	70-130	30	5 5 3
Cadmium	6020	70-130	30	1
Calcium	6010	70-130	30	50
Chromium	6010	70-130	30	3
Chromium	6020	70-130	30	4
Chromium VI	7196	D-130	30	0.2
Cobalt	6010	70-130	30 30	
Cobalt	6020	70-130 70-130	30	3
Copper	6010	70-130	30 30	4 3
F.F	0010	/U-13U	30	3

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TABLE 5-3 Quality Assurance Objectives for Solid Waste / Hazardous Waste Methods

CONSTITUENT	<u>Method</u>	ACCURACY % REC-AR	PRECISION RPD-MAV	DLR mg/L
CONSTITUENT	Methou	70 KEC-AK	ICI D-IVIA V	1112/12
Trace Metals continued				
Copper	6020	70-130	30	4
Gold	231.1	70-130	30	4 3 3 4
Iron	6010	70-130	30	3
Lead	7420	70-130	30	4
Lead	6020	70-130	30	1
Organic Lead	LUFT	50-150	20	4
Lithium	7430	70-130	30	3
Magnesium	6010	70-130	30	50
Manganese	6010	70-130	30	2
Manganese	6020	70-130	30	2 2
Mercury	7471	65-135	30	0.01
Molybdenum	6010	70-130	30	3
Molybdenum	6020	70-130	30	1
Nickel	6010	70-130	30	3 2
Nickel	6020	70-130	30	2
Phosphorous, Total	6010	65-135	30	50
Potassium	6010	70-130	30	50
Selenium	7740	65-135	30	3 4
Selenium	6020	65-135	30	4
Silver	6010	70-130	30	3
Silver	6020	70-130	30	1
Sodium	6010	70-130	30	50
Strontium	6010	65-135	30	3 3 1
Thallium	7841	65-135	30	3
Thallium	6020	65-135	30	
Tin	6010	65-135	30	3
Titanium	6010	65-135	30	5
Uranium	6010	65-135	30	3 5 5 1
Vanadium	6010	70-130	30	1
Zinc	6010	70-130	30	3
Zinc	6020	70-130	30	10

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TABLE 5-3 Quality Assurance Objectives for Solid Waste / Hazardous Waste Methods

CONSTITUENT	Method	ACCURACY <u>% REC-AR</u>	PRECISION <u>RPD-MAV</u>	MDA pCi/Kg
Radiochemistry				
Gamma Emitters	901.1	80-120	20	1
Gross Alpha	900.0	60-140	30	1
Gross Beta	900.0	60-140	30	1
Radon	913.0	N/A	20	10
Strontium 90	905.0	60-140	30	1
Total Radium	900.1	75-125	20	1
Radium 226	903.1	75-125	20	1
Radium 228	904.0	75-125	20	1
Tritium	906.0	75-125	20	300
Uranium	908.0	75-125	20	1

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TABLE 5-4 Quality Assurance Objectives for Ambient Air / Air Pollution Methods

CONSTITUENT	ACCURACY % REC-AR	PRECISION RPD-MAV	DLR ug/m³
Method EPA TO-14			
Surrogates			
2-Bromochloropropane	70-130	N/A	N/A
Analytes			
Acetone	70-130	30	2
Benzene	70-130	30	0.5
Bromodichloromethane	70-130	30	1
Bromoform	70-130	30	ī
Bromomethane	70-130	30	1
1,3-Butadiene	70-130	30	ī
2-Butanone (MEK)	70-130	30	ī
Carbon Disulfide	70-130	30	5
Carbon tetrachloride	70-130	30	1
Chlorobenzene	70-130	30	0.5
Chloroethane	70-130	30	0.5
2-Chloroethyl vinyl ether	70-130	30	
Chloroform	70-130	30	5 3
Chloromethane	70-130	30	0.5
Dibromochloromethane	70-130	30	1
1,2-Dibromoethane (EDB)	70-130	30	2
1,3-Dichlorobenzene	70-130	30	<u>-</u>
1,4-Dichlorobenzene	70-130	30	î
1,2-Dichlorobenzene	70-130	30	î
1,1-Dichloroethane	70-130	30	0.5
1,2-Dichloroethane (EDC)	70-130	30	1
1,1-Dichloroethene	70-130	30	î
cis-1,2-Dichloroethene	70-130	30	î
trans-1,2-Dichloroethene	70-130	30	î
1,2-Dichloropropane	70-130	30	0.5
cis-1,3-Dichloropropene	70-130	30	0.5
trans-1,3-Dichloropropene	70-130	30	0.5
Ethyl Benzene	70-130	30	1
2-Hexanone	70-130	30	0.5
Methylene chloride	70-130	30	5
4-Methyl-2-pentanone (MIBK)	70-130	30	0.5
Styrene	70-130	30	1
1,1,2,2-Tetrachloroethane	70-130	30	1
Tetrachloroethylene (PCE)	70-130	30	1
Toluene	70-130	30	ī
1,1,1-Trichloroethane (TCA)	70-130	30	ī
1,1,2-Trichloroethane	70-130	30	ī
Trichloroethene (TCE)	70-130	30	0.5
Trichlorofluoromethane F-11	70-130	30	1
Trichlorotrifluoroethane F-113	70-130	30	2
Vinyl Acetate	70-130	30	2
Vinyl chloride	70-130	30	0.5
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TABLE 5-4 Quality Assurance Objectives for Ambient Air / Air Pollution Methods

CONSTITUENT	ACCURACY <u>% REC-AR</u>	PRECISION <u>RPD-MAV</u>	DLR ug/m³
Method EPA TO-14 continued Xylenes (Total)	70-130	30	1
Total Petroleum Hydrocarbon	70-130	30	4000
Hydrogen Sulfide (Gold Film)	N/A	30	0.2
CONSTITUENT Method ERA TO 2 (modified)	ACCURACY <u>% REC-AR</u>	PRECISION RPD-MAV	DLR mg/m ³
Method EPA TO-3 (modified) Butanes	70-130	30	20
Ethane	70-130	30	10
Hexanes	70-130	30	40
Methane	70-130	30	10
Pentanes	70-130	30	30
Propane	70-130	30	20
_	ACCURACY	PRECISION	DLR
CONSTITUENT	% REC-AR	<u>RPD-MAV</u>	<u>%</u>
Method ASTM 1945			
Carbon Dioxide	70-130	30	0.1
Carbon Monoxide	70-130	30	0.1
Methane	70-130	30	0.005
Nitrogen	70-130	30	0.02
Oxygen	70-130	30	0.01

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Sampling Procedures

Sample collection and sample handling techniques are important aspects of the overall sample analysis process and have a major impact on the quality and validity of the results. Specific containers and preservatives are used to ensure that sample integrity is not lost through volatility or degradation. In addition, contaminants that are likely to interfere or effect the quality of analytical data must be minimized or eliminated. If a client chooses to collect their own samples, experienced lab staff can brief clients by telephone on the proper methods of sample collection. Detailed procedures to ensure sampling consistency and compliance with method requirements are available. The correct container types, bottle sizes, preservatives, container closures, and holding times for sampling are shown in Table 6-1.

6.1 General Precautions

The result of any analytical determination can be no better than the sample on which it is performed. The objective is to obtain a sample that meets the requirements of the sampling program and manage it in such a way that it does not deteriorate or become contaminated before reaching the laboratory. This objective implies that the relative proportions or concentrations of all pertinent components will be the same in the samples as in the material being sampled, and that the sample will be processed in such a way that no significant changes in composition occur before the tests are made.

A sample may be presented to the laboratory for specific determinations with the collector taking responsibility for its validity. Often the laboratory conducts or prescribes the sampling program which is determined in consultation with the user of the test results. Such consultation is essential to ensure the selection of the appropriate sample and analytical methods that provide a true basis for answering the questions that prompted the sampling.

Before filling, rinse the sample bottle two or three times with the water being collected, unless the bottle contains a preservative. Depending on determinations to be performed, fill container full (most organics determinations) or leave space for aeration, mixing, etc. (microbiological analyses). For samples that will be shipped, preferably leave an air space of about one (1) percent of container capacity to allow for thermal expansion. Special precautions are necessary for samples containing organic compounds and trace metals. Because many constituents may be present at concentrations of micrograms per liter, they may be totally or partially lost if proper sampling and preservation procedures are not followed.

Representative samples of some sources can be obtained only by making composites of samples collected over a period of time or at many different sampling points. The details of collection vary so much with local conditions that no specific recommendations would be universally applicable. Sometimes it is more informative to analyze numerous separate samples instead of one composite so as not to obscure high and low results. Sample carefully to ensure that analytical results represent the actual sample composition. The following are several important factors affecting results:

the presence of suspended matter or turbidity; the method chosen for its removal, and; the physical and chemical changes brought about by storage or aeration.

Particular care is required when processing (grinding, blending, sieving, filtering) samples to be analyzed for trace constituents, especially metals and organic compounds. Some determinations, particularly of lead, can be invalidated by contamination from such processing.

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Sampling Procedures

6.1 General Precautions continued

Treat each sample individually with regard to the substances to be determined, the amount and nature of turbidity present, and other conditions that may influence the results.

It is impractical to give directions covering all conditions. The choice of technique for collecting a homogeneous sample must be left to the analyst's judgment. In general, separate any significant amount of suspended matter by decantation, centrifugation, or an appropriate filtration procedure. Often a slight turbidity can be tolerated if experience shows that it will cause no interference in gravimetric or volumetric tests. Its influence can be corrected in colorimetric tests, where it has potentially the greatest interfering effect. When relevant, state whether or not the sample has been filtered. To measure the total amount of a constituent, do not remove suspended solids, but treat them appropriately.

Make a record of every sample collected and identify every bottle, preferably by attaching an appropriately inscribed tag or label. Record sufficient information to provide positive sample identification at a later date. Include the name of the sample collector, the date, hour, and exact location, the water temperature, and any other data that may be needed for correlation, such as weather conditions, water level, stream flow, post-sampling handling, etc. Provide space on the label for the initials of those assuming sample custody and for the time and date of transfer. Identify sampling points by detailed description, by maps, or with the aid of stakes, buoys, or landmarks in a manner that will permit their identification by other persons without reliance on memory or personal guidance. When sample results are expected to be involved in litigation, it is recommended to use formal "chain-of-custody" procedures which trace sample history from collection to final reporting.

Hot samples collected under pressure should be cooled while still under pressure.

Before collecting samples from distribution systems, flush lines sufficiently to insure that the sample is representative of the supply, taking into account the diameter and length of the pipe to be flushed and the velocity of flow.

Collect samples from wells only after the well has been pumped sufficiently to insure that the sample represents the groundwater source. Sometimes it will be necessary to pump at a specified rate to achieve a characteristic drawdown, if this determines the zones from which the well is supplied. Record pumping rate and drawdown.

When samples are collected from a river or stream, observed results may vary with depth, stream flow, and distance from shore and from one shore to the other. If equipment is available, take an "integrated" sample from top to bottom in the middle of the stream or from side to side at mid-depth.

Lakes and reservoirs are subject to considerable variations from normal causes such as seasonal stratification, rainfall, runoff, and wind. Choose location, depth, and frequency of sampling to reflect local conditions and the purpose of the investigation. Avoid surface scum.

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Sampling Procedures

6.1 General Precautions continued

For certain constituents, sampling location is extremely important. Avoid areas of excessive turbulence because of potential loss of volatile constituents and of potential presence of toxic vapors. Avoid sampling at weirs because such locations tend to favor retrieval of lighter-than-water, immiscible compounds. Generally, collect samples beneath the surface in quiescent areas. If composite samples are required, take care that sample constituents are not lost during compositing because of improper handling of the sample being pooled. For example, casual dumping together of portions rather than addition to the composite through a submerged siphon can cause unnecessary volatilization.

Use only representative samples (or those conforming to a sampling program) for examination. The great variety of conditions under which collections must be made makes it impossible to prescribe a fixed procedure. In general, take into account the tests or analyses to be made and the purpose for which the results are needed.

6.1.2 Field Notebook

The sampler or field investigator should keep a field notebook (preferably bound with pages numbered) to record sample collection procedures, dates, laboratory identification, sample collection location, and the name of the sampler. This is important for later recall or legal challenge.

6.2 Sample Collection

6.2.1 Water Sampling

6.2.1.1 Grab or Catch Samples

Strictly speaking, a sample collected at a particular time and place can represent only the composition of the source at that time and place. However, when a source is known to be fairly constant in composition over a considerable period of time or over substantial distances in all directions, then the sample may be said to represent a longer time period or a larger volume, or both, than the specific point at which it was collected. In such circumstances, some sources may be fairly represented by single grab samples. Examples are some water supplies, some surface waters, and rarely, some wastewater streams. When a source is known to vary with time, grab samples collected at suitable intervals and analyzed separately can document the extent, frequency, and duration of these variations. Choose sampling intervals on the basis of the frequency with which changes may be expected, which may vary from as little as five (5) minutes to as long as one (1) hour or more. Seasonal variations in natural systems may necessitate sampling over months. When the source composition varies in space rather than time, collect samples from appropriate locations.

Use great care in sampling wastewater, sludges, sludge banks, and muds. No definite procedure can be given, but take every possible precaution to obtain a representative sample or one conforming to a sampling program.

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Sampling Procedures

6.2.1.2 Composite Samples

In most cases, the term "composite sample" refers to a mixture of grab samples collected at the same sampling point at different times. Sometimes the term "time-composite" is used to distinguish this type of sample from others. Time-composite samples are most useful for observing average concentrations that will be used, for example, in calculating the loading or the efficiency of a wastewater treatment plant. As an alternative to the separate analysis of a large number of samples and the computation of average and total results, composite samples represent a substantial saving in laboratory effort and expense. For these purposes, a composite sample representing a 24 hour period is considered standard for most determinations. Under certain circumstances, however, a composite sample representing one shift, or a shorter time period, or a complete cycle of a periodic operation, may be preferable. To evaluate the effects of special, variable, or irregular discharges and operations, collect composite samples representing the period durig which such discharges occur.

For determining components or characteristics subject to significant and unavoidable changes on storage, do not use composite samples. Make such determinations on individual samples as soon as possible after collection and preferably at the sampling point. Analyses for all dissolved gases, residual chlorine, soluble sulfide, temperature, and pH are examples of this type of determination. Changes in such components as dissolved oxygen or carbon dioxide, pH, or temperature may produce secondary changes in certain inorganic constituents such as iron, manganese, alkalinity, or hardness. Use time-composite samples only for determining components that can be demonstrated to remain unchanged under the conditions of sample collection and preservation.

Take individual portions in a bottle having a diameter of at least 35 mm at the mouth and a capacity of at least 120 mL. Collect these portions every hour, in some cases every half hour or even every five (5) minutes, and mix at the end of the sampling period or combine in a single bottle as collected. If preservatives are used, add them to the sample bottle initially so that all portions of the composite are preserved as soon as collected. Analysis of individual samples sometimes may be necessary. It is desirable, and often essential, to combine individual samples in volumes proportional to flow. A final sample volume of 2 to 3 L is sufficient for sewage, effluents, and wastes.

Automatic sampling devices are available; however, do not use them unless the sample is preserved as described below. Clean sampling devices, including bottles, daily to eliminate biological growths and other deposits.

6.2.1.3 Integrated Samples

For certain purposes, the information needed is provided best by analyzing mixtures of grab samples collected from different points simultaneously, or as nearly so as possible. Such mixtures sometimes are called integrated samples. An example of the need for such sampling occurs in a river or stream that varies in composition across its width and depth. To evaluate average composition or total loading, use a mixture of samples representing various points in the cross-section, in proportion to their relative flows. The need for integrated samples also may exist if combined treatment is proposed for several separate wastewater streams, the interation of which may have a significant effect on treatability or even on composition. Mathematical prediction of the interactions may be inaccurate or impossible and testing a suitable integrated sample may provide more useful information.

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Sampling Procedures

6.2.1.3 Integrated Samples continued

Both natural and artificial lakes show variations of composition with both depth and horizontal location. However, under many conditions, neither total nor average results are especially significant; local variations are more important. In such cases, examine samples separately rather than integrate them.

Preparation of integrated samples usually requires special equipment to collect a sample from a known depth without contaminating it with overlying water. Knowledge of the volume, movement, and composition of the various parts of the water being sampled usually is required. Therefore, collecting integrated samples is a complicated and specialized process that cannot be described in detail.

6.2.2 Soil Sampling

Soil samples collected from a backhoe, the ground or a soil coring device, should be collected in a thin-walled stainless steel or brass cylinder at least three inches long by one inch in diameter that has been prepared by the laboratory doing the analysis or the project consultant (cylinders can be made to fit inside the preferred split- barrel core sampler). About one inch of soil should be removed from the immediate surface area where the sample is to be taken and the cylinder then pounded in to the soil with a mallet or hammer. No headspace should be present in the cylinder once the sample is collected. When the sample is collected, each end of the cylinder should be covered with teflon tape and then capped with a polyethylene lid, taped and labeled. The sample should be immediately placed in an ice chest and kept cool at 4 C for delivery to the laboratory. Care should be taken throughout, to avoid contamination of both the inside and outside of the cylinder and its contents.

In situations where the above procedure is inappropriate (i.e. semi-solid samples), glass vials with Teflon seal and screw cap should be used.

6.2.3 Special Sampling Considerations

6.2.3.1 Volatile Organics including Organic Lead

When collecting the samples, liquids and solids should be introduced into the vials gently to reduce agitation which might drive off volatile compounds. Liquid samples should be poured into the vial without introducing any air bubbles within the vial as it is being filled. Should bubbling occur as a result of violent pouring, the sample must be poured out and the vial refilled. Each VOA vial should be filled until there is a meniscus over the lip of the vial. The screw-top lid with the septum (Teflon side toward the sample) should then be tightened onto the vial. After tightening the lid, the vial should be inverted and tapped to check for air bubbles. If there are any air bubbles present the sample must be retaken. Two VOA vials should be filled per sample location.

VOA vials for samples with solid or semi-solid (sludges) matrices should be completely filled as best as possible. The vials should be tapped slightly as they are filled to eliminate as much free air space as possible. Two vials should also be filled per sample location.

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6.2.3.1 Volatile Organics including Organic Lead continued

VOA vials should be filled and labeled immediately at the point at which the sample is collected. They should NOT be filled near a running motor or any type of exhaust system because discharged fumes and vapors may contaminate the samples. The two vials from each sampling location should then be sealed in separate plastic bags to prevent cross- contamination between samples particularly if the sampled waste is suspected of containing high levels of volatile organics. (Activated carbon may also be included in the bags to prevent cross-contamination from highly contaminated samples). VOA samples may also be contaminated by diffusion of volatile organics through the septum during shipment and storage. To monitor possible contamination, a trip blank prepared from distilled deionized water should be carried throughout the sampling, storage, and shipping process.

6.2.3.2 Semivolatile Organics including Pesticides and Herbicides

Containers used to collect samples for the determination of semivolatile organic compounds should be soap and water washed followed by methanol (or isopropanol) rinsing. The sample containers should be of glass or Teflon and have screw-top covers with Teflon liners. In situations where Teflon is not available, samples may react with the aluminum foil, causing eventual contamination of the sample. Plastic containers or lids may NOT be used for the storage of samples due to the possibility of sample contaminat ion from the phthalate esters and other hydrocarbons within the plastic. Sample containers should be filled with care so as to prevent any portion of the collected sample coming in contact with the field persons gloves, thus causing contamination. Samples should not be collected or stored in the presence of exhaust fumes. If the sample comes in contact with the an automatic sampler run reagent water through the sampler and use as a field blank.

6.2.3.3 Trace Metals

In the determination of trace metals, containers can introduce either positive or negative errors in the measurement of trace metals by (a) contributing contaminants through leaching or surface desorption, and (b) depleting concentrations through adsorption. Thus the collection and treatment of the sample prior to analysis require particular attention.

6.2.3.4 Radiochemistry

6.2.3.4.1 Radon

Sample by slowly running water into a 2 liter or greater bucket until it overflows for 5 minutes. The water entering the bucket should be as free as possible of bubbles. Fill duplicate 250 mL glass boston round bottles under water, taking care to release all of the air bubbles from inside the bottle. Cap each bottle tightly whil still under water. Dry the bottles and place electrical tape around the cap. Record the date and time. Ship immediately to the laboratory and keep the samples cool at 4 C.

6.2.3.4.1 Radium

The sample should be acidified at the time of sampling.

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6.2.3.5 Air Toxics

In the determination of air toxics there are many ways in which to sample and handle samples. For specifics regarding air toxics sampling and handling please refer to the air toxics sampling standard operating procedure (Document ID S2AIR005.XXX).

6.2.3.6 Product Samples

Free Floating Product (from a well): Sampling of free floating product on the surface of ground water should not be performed until the well has been allowed to stabilize for at least 24 hours after development or other withdrawal procedure. A sample should be collected that is indicative of the thickness of floating product within the monitoring well. This may be accomplished by the use of a clear, acrylic bailer designed to collect a liquid sample where free product and ground water meet. A graduated scale on the bailer is helpful for determining the thickness of free product. Samples should be field-inspected for the presence of odor and/or sheen in addition to the above evaluation. Electronic measuring devices also are available for determining the thickness of the hydrocarbon layer floating on ground water.

6.2.3.7 Aqueous/Dissolved Product

If free product (from a well) is detected, analysis of water for dissolved product chould be conducted after the free product has been substantially removed from the well. Before collecting a water sample a well should be purged until temperature, conductivity and pH stabilize. Often, this will require removal of four or more well volumes by bailing or pumping. Once well volumes are removed and well water is stabilized, a sample can be taken after the water level approaches 80 percent of its initial level. Where water level recovery is slow, the sample can be collected after stabilization is achieved.

Ground water samples should be collected in a manner which reduces or eliminates the possibility of loss of volatile constituents from the sample. For collecting samples, a gas-actuated positive displacement pump or a submersible pump is preferred. A Teflon or stainless steel bailer is acceptable. Peristaltic pumps or airlift pumps should not be used.

Cross-contamination from transferring pumps (or bailers) from well to well can occur and should be avoided by thorough cleaning between sampling episodes. Dedicated (i.e., permanent installation) well pumps, while expensive, are often cost effective in the long term and ensure data reliability relative to cross-contamination. If transfer of equipment is necessary, sampling should proceed from the least contaminated to the most contaminated well, if the latter information is available before sample collection.

Water samples should be collected in vials or containers specifically designed to prevent loss of volatile constituents from the sample. These vials should be provided by an analytical laboratory, and preferably, the laboratory conducting the analysis. No headspace should be present in the sample container once the container has been capped. This can be checked by inverting the bottle, once the sample is collected, and looking for bubbles. Sometimes it is not possible to collect a sample without air bubbles, particularly if water is aerated. In these cases, the investigator should record the problem and account for probable error. Cooling samples may also produce headspace (bubbles), but these will disappear once the sample is warmed for analysis.

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6.2.3.7 Aqueous/Dissolved Product continued

Samples should be placed in an ice chest maintained at 4 C with blue ice (care should be taken to prevent freezing of the water and bursting of the glass vial). A thermometer with a protected bulb should be carried in each ice chest.

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Sampling Procedures

6.3 Sample Handling Policy

Proper sample containers, sample volumes, preservatives, and holding times are essential to providing reliable data. Table 6-1 provides information for each of these items. FGL references the following sources for compiling Table 6-1.

- (1) Federal Register, Volume 49, No. 209, October 26, 1984 and subsequent updates.
- (2) "Handbook for Sampling and Sample Preservation of Water and Wastewater", EPA Method Book, EPA-600/4-82-029, September 1982.
- (3) "Methods for Chemical Analysis in Waters and Waste" (MCAWW) EPA-600/4-79-020
- (4) "Methods for Evaluating Solid Waste", EPA Method Book, SW- 846, rev. 3, and Proposed Revisions.
- (5) "Standard Methods for the Analysis of Water and Wastewater", 17th Edition, 1990.
- (6) "Methods for the Determination of Organic Compounds in Drinking Water", EPA Method Book, EPA-600/4-88-039, December 1988.
- (7) "Methods for the Determination of Organic Compounds in Drinking Water-Supplement I", EPA Method Book, EPA-600/4- 90-020, July 1990.
- (8) "Methods for the Determination of Organic Compounds in Drinking Water-Supplement II", EPA Method Book, EPA-600/4- 90-020, July 1990.
- (9) "Methods for Organic Chemical Analysis of Municipal and Industrial Wastewater", EPA Method Book, EPA 600/4-82-057, July 1982.
- (10) "Prescribed Procedures for Measurement of Radioactivity in Drinking Water", EPA Method Book, EPA-600/4-80-032, August 1980.

6.3.1 Container Check Policy

Sample bottles for most analyses, such as metals, and organics analyses, are purchased precleaned according to EPA Protocol specification from various vendors. Cases of sample bottles are logged in upon receipt. The log book contains the bottle type, lot number or manufacture date, receive date and number of cases. This information is also recorded on the document provided by the manufacturer and is retained in a file for that bottle type. Most containers are checked and documented by the manufacturer. Bacteriology and Santa Paula metals bottles must be checked in-house by lot or on a manufacture date basis. If neither of these are available then one container must be checked from every ten cases. The following files are to be maintained for storing certificates or in-house analysis data:

```
40 mL VOA - for all VOA styles
125 mL Boston Round (B.R.) - for all 125 mL B.R. bottle styles
250 mL Boston Round (B.R.) - for all 250 mL B.R. bottle styles
1 L Boston Round (B.R.)
250 mL Wide Mouth (W.M.)
1 Qt. Wide Mouth (W.M.)
4 oz. Bact
500 mL Plastic (SP only, for Al, Sb, As, Ba, Be, Cd, Ca, Cr, Co, Cu, Fe, Pb, Mg, Mn, Hg, Ni, K, Se, Ag, Na, Tl, V, Zn)
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For those bottles not listed and not verified (such as 1 Qt plastic) it has been determined that the tests performed from those containers are not at risk from contamination by the container.

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Sampling Procedures

6.3.1 Container Check Policy continued

When an in-house verification is required the sample is given a lab number, chain-of-custody and is treated as a regular sample. The results are formally reported but only the signature of the QA Director or Officer to verify cleanliness is required prior to filing.

Lots determined to be contaminated are returned and replaced.

6.3.2 Shipping Samples

Prior to shipment of samples, all documentation must be ready for proper chain of custody. The information necessary for documenting chain of custody is outlined in the following section of the quality assurance manual (section 7). After filling out the proper sample documentation, the samples and documents should be placed in an ice chest with adequate protection. Normally "Blue Ice" is used for keeping samples cool. However, dry ice may be used if approved by Department of Transportation (DOT).

6.3.3 Sample Kits

FGL Environmental supplies the appropriate sample containers, preservatives, chain-of-custody forms, coolers with blue ice, and packing materials to client upon request. There is no charge for these services as long as FGL is the laboratory receiving the samples for analysis. Arrangements for sample kits may be made through the client services department.

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Sampling Procedures

TABLE 6-1 RECOMMENDED SAMPLE COLLECTION AND PRESERVATION

<u>Analysis</u>	Container	Volume (mL)	Preservation	Holding <u>Time</u>			
General Inorganic Chemistry							
Acidity	P,G	250	Cool, 4 C	14 days			
Alkalinity	P,G	250	Cool, 4 C	14 days			
Asbestos	G	1000	Cool, 4 C	48 hr.			
Bicarbonate	P	250	Cool, 4 C	14 days			
Biochemical Oxygen Demand	P,G	1000	Cool, 4 C	48 hr.			
Boron	P	250	Cool, 4 C	6 mo.			
Bromide	P	250	Cool, 4 C	28 days			
Carbonate	P	250	Cool, 4 C	14 days			
Carbon Dioxide	P,G	250	Cool, 4 C	immed.			
Chemical Oxygen Demand	P,G	250	H ₂ SO ₄ , pH<2; Cool, 4 C	28 days			
Chloride	P,G	250	Cool, 4 C	28 days			
Chlorine Residual	P,G	500	Cool, 4 C	2 hr.			
Chlorine Demand	P,G	2000	Cool, 4 C	2 hr.			
Color	P,G	250	Cool, 4 C	48 hr.			
Cyanide, Total	P,G	1000	NaOH, pH>12; Cool, 4 C	14 days			
Electrical Conductivity	P	250	Cool, 4 C	28 days			
Fluoride	P,G	250	Cool, 4 C	28 days			
Hardness, Total	P,G	250	HNO ₃ , pH<2; Cool, 4 C	6 mo.			
Hydroxide	P,G	250	Cool, 4 C	14 days			
Iodide	P,G	250	Cool, 4 C	24 hr.			
Langelier Index	P,G	500	Cool, 4 C	2 hr.			
MBĀS	P,G	500	Cool, 4 C	48 hr.			
Nitrogen,	•		,				
Ammonia	P,G	250	H ₂ SO ₄ , pH<2; Cool, 4 C	28 days			
Nitrate+Nitrite	P,G	250	H ₂ SO ₄ , pH<2; Cool, 4 C	28 days			
Nitrate	P,G	250	Cool, 4 C	48 hr.			
Nitrite	P,G	250	Cool, 4 C	48 hr.			
Organic	P,G	400	H ₂ SO ₄ , pH<2; Cool, 4 C	28 days			
Total	P,G	250	H ₂ SO ₂ , pH<2; Cool, 4 C	28 days			
Total Kjeldahl	P,G	250	H ₂ SO ₄ , pH<2; Cool, 4 C	28 days			
Odor	G	500	Cool, 4 C	24 hr.			
Oil and Grease	G	1000	H ₂ SO ₄ , pH<2; Cool, 4 C	28 days			
Oxygen, Dissolved	G	250	Cool, 4 C	immed.			
w/glass stopper							
pН	P,G	250	Cool, 4 C	2 hr.			
Phenolics	G	500	H_2SO_4 , pH <2; Cool, 4 C	28 days			

P = plastic, G = glass Note: All solid samples should be collected in stainless steel sleeves, brass sleeves or in glass jars all with teflon-lined caps and 4-8 oz. capacity. All solid samples should be kept cool at 4 C.

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Sampling Procedures

TABLE 6-1 RECOMMENDED SAMPLE COLLECTION AND PRESERVATION

<u>Analysis</u>	Container	Volume (mL)	Preservation	Holding <u>Time</u>
•		(1112)	Teservation	Time
General Inorganic Chemisti	y continued			
Phosphorus				
Ortho/dissolved	P,G	250	Cool, 4 C	48 hr.
Total	P,G	250	H ₂ SO ₄ , pH<2; Cool, 4 C	28 days
Resistivity	P	250	Cool, 4 C	28 days
Silica	P	250	Cool, 4 C	28 days
Sodium Absorption Ratio	P	250	HNO3, pH<2	6 mo.
Solids,				
Filterable	P,G	250	Cool, 4 C	7 days
Non-filterable	P,G	250	Cool, 4 C	7 days
Total	P,G	250	Cool, 4 C	7 days
Volatile	P,G	250	Cool, 4 C	7 days
Settleable	P,G	1000	Cool, 4 C	48 hr.
Sulfate	P,G	250	Cool, 4 C	28 days
Sulfide				
Total	P,G	500	2 ml ZnAcetate+NaOH, pH>9	7 days
Dissolved	P,G	500	NaOH, pH>9	7 days
Tannin & Lignin	G	250		
Titration - pH adjustment	P,G	250	Cool, 4 C	14 days
Turbidity	P,G	250	Cool, 4 C	48 hr.
Trace Metals				
Chromium VI	P,G	500	Cool, 4 C	24 hr.
Mercury	P,G	500	HNO ₃ , pH<2	28 days
All other metals	P	500	HNO_3 , $pH<2$	6 mo.
			J	
Radiochemistry	D.	1000	IINO II a	(
Gross Alpha & Beta**	P	1000	HNO ₃ , pH <2	6 mo.
Total Radium	P	1000	HNO ₃ , pH <2	6 mo.
Total Uranium	P	1000	HCl, pH <2	6 mo.
Radon*	G	2x250	Cool, 4 C	4 days 6 mo.
Tritium	G P	2x250	Cool, 4 C	6 mo.
Strontium 90	r	1000	HCl, pH <2	o mo.

P = plastic, G = glass

^{*} No headspace over sample

^{**} For non-preserved samples, the holding time is 5 days. For preserved samples, please provide either a non-preserved sample (100 mL) or the electrical conductivity prior to acidification.

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Sampling Procedures

TABLE 6-1 RECOMMENDED SAMPLE COLLECTION AND PRESERVATION

Analysis	Containe	Volume er (mL)	Preservation	Holding <u>Time</u>	
All Bacteriological	P,G	100	0.008% Na ₂ S ₂ 0 ₃ ; Cool, 4 C, Sterile	30 hr.	
Analysis Organic Chemicals	<u>Container</u>	Volume (mL)	Preservation	Holding <u>Time</u>	
Drinking Water					
EPA 501*	G, VOA TFE-septa cap	2 x 40	$Na_2S_2O_3$, if chlorinated HCl, pH <2; Cool, 4 C	14 days	
EPA 502.2*	G, VOA TFE-septa cap	2 x 40	Na ₂ S ₂ O ₃ , if chlorinated HCl pH <2; Cool, 4 C	14 days	
EPA 504*	G TFE-septa cap	2 x 40	Na ₂ S ₂ O ₃ , if chlorinated Cool, 4 C	28 days	
EPA 505**	G TFE-septa cap	2 x 40	Na ₂ S ₂ O ₃ , if chlorinated Cool, 4 C	7 days	
EPA 507	G, amber TFE-lined cap	1 x 1000	Na ₂ S ₂ O ₃ , if chlorinated or HCl pH <2; Cool, 4 C	14 days	
EPA 508**	G, amber TFE-lined cap	1 x 1000	Na ₂ S ₂ O ₃ , if chlorinated Cool, 4 C	7 days	
EPA 510*	G, amber TFE-septa cap	1 x 250	Na ₂ S ₂ O ₃ , if chlorinated Cool, 4 C	14 days	
EPA 515.1**	G, amber TFE-lined cap	1 x 1000	Na ₂ S ₂ O ₃ , if chlorinated Cool, 4 C	7 days	
EPA 524.2*	G, VOA TFE-septa cap	2 x 40	Na ₂ S ₂ O ₃ , if chlorinated or HCl pH <2; Cool, 4 C	14 days	
EPA 525**	G, amber TFE-lined cap	1 x 1000	Cool, 4 C	7 days	

^{*} No head space over sample.

** This is the maximum holding time prior to extraction. The extracted sample may be held up to 40 days before analysis.

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Sampling Procedures

TABLE 6-1 RECOMMENDED SAMPLE COLLECTION AND PRESERVATION

<u>Analysis</u>	<u>Container</u>	Volume (mL)	Preservation	Holding <u>Time</u>
Organic Chemicals				
Drinking Water				
EPA 531	G, amber	1 x 250	Na ₂ S ₂ O ₃ , if chlorinated Monochloroacetic acid buffer	14 days
EPA 547	G, amber	1 x 125	Na ₂ S ₂ O ₃ , if chlorinated Cool, 4 C	6 mo.
EPA 548	48 G, amber		Na ₂ S ₂ O ₃ , if chlorinated Cool, 4 C	7 days
EPA 549	G, amber silanized	1 x 1000	Na ₂ S ₂ O ₃ , if chlorinated Cool, 4 C	7 days
EPA 550.1	G, amber	1 x 1000	Na ₂ S ₂ O ₃ , if chlorinated Cool, 4 C	7 days
EPA 551	G, amber	2 x 40	NH ₄ Cl, Cool, 4 C	28 days
EPA 552	G, amber	1 x 1000	NH ₄ Cl, Cool, 4 C	14 days
EPA 1613A	G, amber	1 x 1000	Na ₂ S ₂ O ₃ , if chlorinated Cool, 4 C	7 days
Wastewater and Ha	zardous Waste			
EPA 601/8010*	G, VOA TFE-septa cap	2 x 40	Na ₂ S ₂ O ₃ , if chlorinated HCl, pH <2; Cool, 4 C	14 days
EPA 8011	G, VOA TFE-septa cap	2 x 40	Na ₂ S ₂ O ₃ , if chlorinated Cool, 4 C	28 days
EPA 602/8020*	G, VOA TFE-septa cap	2 x 40	Na ₂ S ₂ O ₃ , if chlorinated HCl, pH <2; Cool, 4 C	14 days
EPA 603/8030*	G, VOA TFE-septa cap	2 x 40	Na ₂ S ₂ O ₃ , if chlorinated Adjust pH to 4-5; Cool, 4 C	14 days
EPA 604/8040**	G, amber TFE-lined cap	1 x 1000	Na ₂ S ₂ O ₃ , if chlorinated Cool, 4 C	7 days

^{*} No head space over sample.

** This is the maximum holding time prior to extraction. The extracted sample may be held to 40 days before analysis.

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Sampling Procedures

TABLE 6-1 RECOMMENDED SAMPLE COLLECTION AND PRESERVATION

<u>Analysis</u>	Container	Volume (mL)	Preservation	Holding <u>Time</u>
Organic Chemicals				
Wastewater and Ha	zardous Waste			
EPA 608/8080**	G, amber TFE-lined cap	1 x 1000	Na ₂ S ₂ O ₃ , if chlorinated Cool, 4 C	7 days
EPA 610/8310	CPA 610/8310 G, amber TFE-lined cap		Na ₂ S ₂ O ₃ , if chlorinated Cool, 4 C	14 days
EPA 613**	G, amber TFE-lined cap	1 x 1000	Na ₂ S ₂ O ₃ , if chlorinated Cool, 4 C	7 days
EPA 614/8140**	G, amber TFE-lined cap	1 x 1000	Cool, 4 C	7 days
EPA 615/8150**	G, amber TFE-lined cap	1 x 1000	Na ₂ S ₂ O ₃ , if chlorinated Cool, 4 C	7 days
EPA 619	G, amber TFE-lined cap	1 x 1000	Na ₂ S ₂ O ₃ , if chlorinated Cool, 4 C	14 days
EPA 624/8240*	G, VOA TFE-septa cap	2 x 40	Na ₂ S ₂ O ₃ , if chlorinated HCl, pH <2; Cool, 4 C	14 days
EPA 625/8270**	G, amber TFE-lined cap	1 x 1000	Na ₂ S ₂ O ₃ , if chlorinated Cool, 4 C	7 days
EPA 632**	G, amber TFE-lined cap	1 x 1000	Cool, 4 C	7 days
EPA 9020* (TOX)***	G, amber TFE-lined cap	1 x 250	H ₂ SO ₄ , pH <2 Cool, 4 C	28 days*
EPA 415.1/9060 (TOC)	G, amber TFE-lined cap	1 x 250	HCl or H ₂ SO ₄ , pH <2 Cool, 4 C	28 days
Tributyltin	G, amber TFE-lined cap	2 x 1000	Cool, 4 C	28 days

^{*} No head space over sample.

^{**} This is the maximum holding time prior to extraction. The extracted sample may be held up to 40 days before analysis.

^{***} RCRA holding time is 7 days.

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Sampling Procedures

TABLE 6-1 RECOMMENDED SAMPLE COLLECTION AND PRESERVATION

<u>Analysis</u>	<u>Container</u>	Volume (mL)	Preservation	Holding <u>Time</u>				
Organic Chemicals								
Underground Storag	ge Tank Analyses							
EPA 418.1	G, amber TFE-lined cap	1 x 1000	Cool, 4 C	28 days				
EPA 8015,8015M*	G, VOA TFE-septa cap	2 x 40	HCl, pH <2 Cool, 4 C	14 days				
EPA 601/8010*	G, VOA TFE-septa cap	2 x 40	Na ₂ S ₂ O ₃ , if chlorinated HCl, pH <2; Cool, 4 C	14 days				
EPA 602/8020*	G, VOA TFE-septa cap	2 x 40	Na ₂ S ₂ O ₃ , if chlorinated HCl, pH <2; Cool, 4 C	14 days				
Hazardous Waste C	haracterization							
Corrosivity	P,G	100	Cool, 4 C	7 days				
Ignitability	G TFE-lined cap	100	Cool, 4 C	7 days				
Reactivity/ Reactions	G TFE-lined cap	100	Cool, 4 C	7 days				
Sulfide/Sulfide generation	G TFE-lined cap	100	Cool, 4 C	7 days				

^{*} No head space over sample.

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Sampling Procedures

TABLE 6-1 RECOMMENDED SAMPLE COLLECTION AND PRESERVATION

<u>Analysis</u>	<u>Container</u>	Volume (mL)	Preservation	Holding <u>Time</u>
TTLC, STLC, TCL	P, and EP Toxicit	y		
Metals	G TFE-lined cap	500	Cool, 4 C	30 days
Pesticides	G, amber TFE-lined cap	1000	Cool, 4 C	7 days
Herbicides	G, amber TFE-lined cap	1000	Cool, 4 C	7 days
Bioassays				
Chronic	P,G	3x1000	Cool, 4 C	24 hr.
Acute	P,G	2x5 gal	Cool, 4 C	24 hr.

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Sample Custody

7.1 Sample Custody

It is essential to ensure sample integrity from the time of collection through analysis and final disposition. This includes the ability to trace possession and handling of the samples. This is referred to as chain-of-custody and is important in the event of litigation involving the results. Where litigation is not involved, chain-of-custody procedures are useful for routine control of sample flow.

A sample is considered to be under a person's custody if it is in the individual's physical possession, in the individual's sight, secured in a tamper-proof manner by that individual, or is secured in an area restricted to authorized personnel. The following procedures summarize the major aspects of chain-of-custody.

7.1.1 Sample Labels

Use labels to prevent sample misidentification. Gummed paper labels or tags generally are adequate. Include at least the following information: sample number, name of collector, date and time of collection, and place of collection. Affix labels to sample containers before or at the time of sampling. Fill label out with waterproof ink at time of collection.

7.1.2 Custody Seals

Use sample seals to detect unauthorized tampering with samples up to the time of analysis. Plastic seals are normally used. Attach seal in such a way that it is necessary to break the seal to open the sample container. Affix seal to container before sample leaves custody of sampling personnel.

7.1.3 Field Log Book

Record all information pertinent to a field survey or sampling in a bound log book. As a minimum, include the following in the log book: purpose of sampling; location of sampling point; name and address of field contact; producer of material being sampled and address, if different from location; and type of sample. Because sampling situations vary widely, no general rule can be given as to the information to be entered in the log book. It is desirable to record sufficient information between the logbook and chain-of-custody so that one could reconstruct the sampling without reliance on the collector's memory. Protect the log book and keep it in a safe place.

7.1.4 Shipping Samples or Sample Delivery to Laboratory

Prior to shipping samples all documentation must be ready for proper chain of custody. The information necessary for documenting chain of custody is outlined section 7.3. After filling out the proper sample documentation, the samples and documents should be placed in an ice chest with adequate protection. Normally "Blue Ice" is used for keeping samples cool. However, dry ice may be used if approved by Department of Transportation (DOT).

If client provides direct delivery of sample to laboratory, samples should be delivered as soon as practical. Documentation must be ready for proper chain of custody. Again, all information necessary for documenting chain of custody is outlined section 7.3. Accompany sample with chain- of-custody record and a sample analysis request sheet. Deliver sample to sample custodian.

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Sample Custody

7.1.6 Receipt and Logging of Sample

In the laboratory, the sample custodian receives the sample and inspects its condition and seal, reconciles label information and seal against the chain-of-custody record, assigns a laboratory number, logs sample in the laboratory computer, and stores it in a secured storage room or cabinet until it is assigned to an analyst.

7.1.7 Assignment of Sample for Analysis

The laboratory supervisor usually assigns the sample for analysis. Once in the laboratory, the supervisor or analyst is responsible for the sample's care and custody.

7.1.8 Safety Considerations

Because sample constituents can be toxic, take adequate precautions during sampling and sample handling. Toxic substances can enter through the skin and, in the case of vapors, through the lungs. Inadvertent ingestion can occur via direct contact with foods or by adsorption of vapors onto foods. Precautions may be limited to wearing gloves or may include coveralls, aprons, or other protective apparel. Always wear eye protection. When toxic vapors might be present, sample only in well-ventilated areas or use a respirator or self-contained breathing apparatus. In a laboratory, open sample containers in a fume hood. Never have food near samples or sampling locations; always wash hands thoroughly before handling food.

If there is any possibility that flammable organic compounds may be present, take adequate precautions. Prohibit smoking near samples, sampling locations, and in the laboratory. Keep sparks, flames, and excessive heat sources away from samples, and sampling locations.

Radioactivity is screened at the time of sample receipt. Consult the radiation safety SOP and Radiation Safety Officer for proper handling of samples.

7.2 Laboratory Sample Control and Tracking

FGL's sample control objectives are achieved through the use of a Laboratory Information Management System (LIMS). LIMS is a computer software system specifically designed by FGL for tracking and handling of the large amount of information required to efficiently manage an analytical chemistry laboratory. The system provides a versatile, easy-to-use vehicle for the laboratory managers and chemists to perform sample tracking and status checks.

7.3 Sample Receiving Policy

Obtain a chain-of-custody record accompanying each sample or group of samples. The chain-of-custody is usually prepared in the field.

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Sample Custody

7.3.1 Obtain the following client information and record on chain of custody:	
Reporting -	
Address	
Phone Number	
Fax Number	
Person to Contact	
Billing -	
Address	
Phone Number	
Fax Number	
P.O. or contract Number	
Person to Contact	
7.3.2 Obtain the following project and sample information and record on chain of custody:	
Project description	
Sample descriptions	
Sample type	
Sampling date and time	
Sample containers, preservatives	
EPA Method Numbers or method descriptions	
Report form required: State FGL	
Determine turn-around-time requirement:	
RushNumber of Days	
7.3.3 Inspect the sample for the following:	
Have holding times been observed and determine if it is possible for FGL to meet	
holding times?	
Has the correct preservative been used?	
Is the sample size adequate?	
Is the sample container satisfactory?	
Note sample condition:	
Broken/leaking container	
TemperatureAmbientChilled	
Record Actual Temperature	
Check for headspace when appropriate	
Also note on the chain of custody any problems with sample condition, the person notified time and date notified, and customers response, if any.	d
Saven all samples for radio showing barand using the Cairon sounts when in the sample	
Screen all samples for radio chemical hazard using the Geiger counter kept in the sample receiving area. Consult the radiation safety SOP and Radiation Safety Officer for prope	
handling of samples.	
7.3.4 Log the sample information into the LIMS under one of the following divisions:	
Inorganic	
Organic	
Radiochemistry	
Bacteriology	
Agronomics	

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Sample Custody

7.3.5 Transfer samples and analyses worksheets to the proper refrigerator or lab work distribution area.

7.4 Sample Storage

All samples are retained for a minimum of 30 days except for microbiological samples which are held for a shorter time. A longer storage period can be arranged at the request of the client.

7.5 Sample Disposal

All samples which are considered to be potentially hazardous based upon analytical results or matrix, will be disposed of by lab packing. Extremely hazardous samples may be returned to the client for disposal. All disposal arrangements should be made with a project manager.

7.6 Subcontracted Lab Work

On occasion, laboratory work may need to be subcontracted to certified labs approved by FGL Environmental. Prior to subcontracting the client will be notified. Under no circumstances will work be subcontracted without client requirements being met. When work is subcontracted, it is done so under chain of custody, and the proper records are included with the data package. When work is subcontracted between the FGL laboratories the original chain of custody is carried forward with the samples. When work is subcontracted to an outside laboratory a separate chain of custody is prepared.

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Sample Custody

Figure 7-1 Chain of Custody

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CHAIN OF CUSTODY AND ANALYSIS REQUEST DOCUMENT

Information and directions for completion

The attached document (Chain of Custody) is designed to encompass all the pertinent information pertaining to the sample(s) to be collected/received and the subsequent analysis(es).

Please note that all sampling and analyses conducted by this laboratory are subject to the terms and conditions on the reverse side of the Chain of Custody.

SECTION I This section should be fully completed and clearly printed. If the billing address differs from the regular address, this should be especially noted. For routine sampling, please indicate if pre log program is required.

SECTION II If the sample is to be taken by you, please clearly print the name of the sampler, the date and time. If the sample is to be taken by FGL, our certificated sampler will be responsible for this information.

SECTION III Please indicate if a rush analysis is requested. If so, please circle the appropriate rush period required. When the chain is received at the lab, you will be notified if this rush is not available. Please indicate if a QA/QC report or a state form is required. The laboratory number and location will be assigned by FGL when the sample(s) are received at the laboratory.

SECTION IV Please number by sample and not by individual test. Each test required (from the numbered sample), should be entered in the diagonal spaces provided to the right. In addition, please ensure that all information is entered in the appropriate boxes, on the same line as the numbered sample.

SECTION V Please use this section to make specific notes; i.e. special sampling, reporting instructions, etc.

SECTION VI requires a signature by a person authorized to relinquish the sample to an FGL representative. If a sample is being shipped, a relinquishing signature is required prior to releasing the sample (properly packaged) to the shipping agent. The receipt of a copy of the shipping document, maintains the integrity of the Chain of Custody.

The correct container(s) to be used for individual tests, with appropriate preservatives can be obtained from FGL. When calling the laboratory for containers, please be specific about the analyses required and if applicable, the number of locations. Samples must be kept cool. Coolers will be provided if requested.

Color Coding of Containers All containers which contain a preservative are marked with a Color Printed Label. The color scheme conforms to industry standards where applicable, and is provided as a quick reference for locating the correct container. Preservative and corresponding labels colors are set out hereunder:

Blue	HCL pH < 2	Black	Monochloroacetic Buffer
Gold	H2SO4 pH < 2	Green	NaOH pH > 12 or NaOH + Zinc Acetate
Red	HNO3 pH < 2	Black	Na2S2O3
	•	White	Other

WE ARE PLEASED TO HAVE THIS OPPORTUNITY TO SERVE YOU

If you have any questions or need sampling assistance, do not hesitate to call us at:

Santa Paula (805) 659 0910 Stockton (209) 942 0181



Chain of CusiOL I AND ANALYSIS REQUEST DOCUMENT

ORIGINAL

CLIENT	DETAILS	1.00	··· SECTIO	NI	SA	MI	PL	NG					S	ECTI	ON II		REP(ORT	INF	ORN	IATIO)N	SECTI	ON III
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	Cus	tomer Number:			1												A/QC r	eport re	quired	l: yes			no	
Address:				_												Ε	lectroni	c Data	Transfe	er: yes			no	
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FAX:				_ [110	ne: _				M	licag	e:												
Project name:					١											┛	ab Loc:	SP		STI	·	VI 🔾		
Contact person	i:						8					_					-	7	7			77	7	
Billing Informa	ation (if different from above)				ı		1					ace (RPL)	ខ្ល	.				/	I	'				
Name:					1		X					place	Produce	HC HC					' /					ace
					_©		Type of Containers: (G) Glass (P) Plastic (V) VOA (MT) Metal Tube		در در	Oil	(W) Waste	SPL) Repl	PRD) PA	rreservauve: (1) naOH + LnAC, (2) naOH, (3) HC! (4) H2SO4. (5) HNO3. (6) Na2S203. (7) Other										OA Headspace
Phone:					Grab(G)		3		Monitoring We Travel Blank Drinking Water	(O) P		Special (SPL)	ssue (2) NaOH, 3, (7) Othe										§
FAX:					Q		last.		Mon: Trave Drink	Sol	Sour	E S	E I	Ac, (2S20									rature (HS)	
Contact person:	:				Composite(C)		Ē	Non-Potable	(MM) (E) (W) (D)	SLD	<u>ي</u>	<u>a</u>	Petio	+ Ch		8		1	1	/ /		11	en (
Purchase order/	contract/FGL quote number:			_	l mo	Z.	Se Se Se Se Se Se Se Se Se Se Se Se Se S	P-Pc	S C 8	lge (S) n	Se pe		3, (6)		<u>s</u>		1 1				/ / 4	n. Iemper Broken	\ <u>&</u>
Pre Log Require	ed: yes no					Containers	;; (G)	(NP) No	Waler Waler Ier) Sluc	Syster	(TOO)	ue (P)	HNO		<u>ğ</u> /						କ୍ରିଞ୍ଚ	<u>e</u>	seal (Y) (N)
AMPLE I	NFORMATION	•	SECTIO	NIV	of Sampling:	r of Cc	Contain	Potable (N	Surface Water Ground Water Wastewater	(SEC	(Sys) System (SRC) Source	Routine (ROUT) Repeat (RPT)	af Tiss	anve: 04, (5)	Vere	3						Sample Condition: T.	Leaking.) sea
umple umber	Location/Descrip	otion	Date Sampled	Time Sampled	8	Number of	Type of	(P) Pot	(SW)	(S) Soil (SLG) Sludge (SLD) Solid (O)	BacT:	BacT: R	(LT) Leaf Tissue (PET) Petiole Tissue	(4) H2SO4, (5) HNO3, (6) Na2S203,	ANAL	A KEQUESTED		\perp	_			Samp	(L) Leak	
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CORPORATE OFFICE & LABORATORY P.O. Box 272/853 Corporation Street Santa Paula, C.A 93061-0272 Tel: (805) 559-0910 FAX: (805) 525-4172 OFFICE & LABORATORY 2500 Stagecoach Road Stockton, CA 95215 Tel: (209) 942-0181 FAX: (209) 942-0423 FIELD OFFICE Visalia, California Tel: (209) 734-9473 Mobile: (209) 737-2399 FAX (200) 734-8435

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Calibration Procedures and Frequency

The production of analytical data of known, defensible and documented quality requires adherence to standardized procedures which cover all aspects of laboratory operation. The following sections provide details of the standardized procedures relating to instrumentation calibration.

8.1 Instrument Calibration

Prior to use, every instrument must be calibrated according to the procedure found in the method specific Standard Operating Procedure (SOP). A list of all laboratory equipment may be found in section 17. Tables 8-1 and 8-2 list the organic mass spectrometer calibration ion abundance criteria which must be met.

The analytical balances are certified once a year by a certified specialist. All balances are labelled as certified. All balances are checked monthly using NIST traceable S weights which are calibrated annually.

All refrigerator, oven, and incubator temperatures are monitored daily, and all oven and incubator thermometers are checked for accuracy on a quarterly basis.

8.2 Calibration Standards

All chemicals used by FGL Environmental are ACS reagent grade, or better. Wherever possible, standards are from sources that are traceable to the National Institude of Standards and Technology. A log book is maintained for all working standards. Each log contains the standard ID or code, date of working standard preparation, analyst initials who prepared the standard, the manufacturers lot number, stock solution expiration date, stock analyte concentration(s), analyte concentration(s), and initial and final volumes used in dilutions. The working standard container is also labeled with ID or code, date of standard preparation, analyst initials who prepared the standard, expiration date, analytes and analyte concentration. The standards are tracked from the logbbok to the analysis in the following manner:

- 1) Working standards and spiking concentrates are labeled and coded. Stock standards should also be labeled and coded if they are used directly for calibration or spiking purposes.
- 2) The standard code is the date the standard was made, the analyst's initials, and the number referring to the first, second, third etc. standard of the day (i.e. 020395KW1).
- 3) The <u>calibration and spiking</u> standard codes should all be recorded with the analytical results. This can be done in several fashions:
 - a. If the test is performed manually the codes should be recorded on the batch sheet.
 - b. If the test is automated the codes can usually be recorded as part of the sample description which is printed by the instrument..
 - c. If the test is calibrated periodically then the original calibration information should have the codes recorded with it (i.e. cyanide, IC, 418.1). The daily calibration CCV's and spike codes should be recorded with each run.

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Calibration Procedures and Frequency

8.3 Calibration Policy

8.3.1 Applicability

This policy is designed to be a guideline to ensure that all data are treated alike, and thus ensuring that data generated on any particular day of analysis are representative of the norm. However, the policies are not intended to be absolute criteria for the acceptance or rejection of any analytical data. In cases where the acceptance criteria outlined in policy or SOPs cannot be achieved then the analyst uses a non-conformance report form to document the difficulty. More than one continuing difficulty will result in a Corrective Action Report (CAR, see section 15). These are on record and will be included in a project data package if that is required by the project plan. An example of a CAR form is shown in figure 15-1.

8.3.2 Linearity

All calibrations should be linear unless otherwise defined in the specific SOP. FGL's definition of linearity is a calibration curve that has a linear regression equal to or greater than 0.995. For organics, if a linear regression is used, a single average response factor may be used if acceptance criteria pass. Specific protocols outlined in a given SOP will take precedence over these generic policies.

8.3.3 Selection of Quantitation Technique (Organics)

For organic analysis, a decision must be made during the validation process (and detailed in the SOP) as to whether an internal or external quantitation technique will be routinely employed.

The internal standard method of quantitation cannot be employed unless all of the following conditions are met:

- (1) The internal standard must be added post-extraction.
- (2) The internal standard must be added quantitatively.
- (3) Any analyte that is a target analyte in the method of interest may not be selected for use as an internal standard.
- (4) The concentration of the internal standard(s) must not exceed the calibration range of the method target analyte. In cases where the target analytes are associated with more than one calibration range (i.e. analytes "1-4" are calibrated from 1 to 10 ppb, while analyte "5" is calibrated from 10 to 100 ppb, all target analytes should be prepared at a level between the highest and lowest calibration standard (e.g. approximately 50 ppb in the example given).

The use of internal standard quantitation is of greatest benefit in those methods subject to injection variability, and thus, variability in the absolute mass injected onto the column(s) employed. The drawback to this technique, for GC methods, is that any compound which exhibits a similar retention time as the compound used for the internal standard will be identified as the internal standard, leading to erronous quantitation. For this reason, the internal standard technique is most useful for GC/MS where deuterated analytes not naturally occuring can be detected and quantified.

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Calibration Procedures and Frequency

8.3.4 Selection of Calibration Method

As part of the validation process, the specific calibration range and calibration method must be determined and documented in the SOP. Once determined in this manner, the same protocols must be followed each time the method is employed. This will ensure that data reduction is not performed differently on separate data sets by different analysts. The calibration acceptance criteria are listed in section 8.3.8. A least squares (linear) regression is initially tried as a calibration method. For organics, if a linear regression is used, a single average response factor may be used. For inorganics, if the acceptance criteria cannot be met using a linear regression, then a second order polynomial can be used to fit the data, with the same acceptance criteria being applied. In the event that neither a simple linear regression nor a second order polynomial fit result in an equation which meets the calibration acceptance criteria, then the calibration range must be reduced.

8.3.5 Minimum Number of Calibration Levels

Most calibrations include a minimum of three or five initial calibration standards plus a blank. Specific SOPs may have other requirements.

8.3.6 Selection of Calibration Levels

Two standards should be included per order of magnitude of concentration of the calibration curve. For example 0.1, 1.0, 10.0 has 2 standards per order magnitude (0.1 and 1.0; 1.0 and 10.0). In cases where instrumentation spans several orders of magnitude, the SOP for that method may not require this policy. The lowest calibration level should be within a factor of 10 of the detection limit for reporting (DLR) for each target compound unless otherwise specified in the SOP.

8.3.7 Calibration Analysis Sequence

The calibration must progress from the analysis of the lowest to highest standard unless the instrumentation does not permit it. A blank must be analyzed after the highest calibration standard.

If the analysis requires an initial high standard to set the gain, a blank must be run before starting with the low calibration standard unless the instrumentation does not permit it.

8.3.8 Calibration Acceptance Criteria

In general, for inorganics, the calculated value for standards must be within 10% of the expected value. However, the value determined by the calibration curve for the lowest standard must be within +/- 50% of the true value and if the calibration is linear through the origin (at less than +/- 1/2 the detection limit). For organics, if a linear regression is used, a single average response factor may be used. The percent relative standard deviation for the individual standard response factors must be less than the maximum value listed for the method in the SOP.

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Calibration Procedures and Frequency

8.3.9 Calibration Check Compounds (CCC) and Initial Calibration Verifications (ICV's)

The CCC or ICV is used to check the validity of the initial calibration. This standard is composed of some or all of the same analytes used for calibration but from a different source than the calibration standard. The standard should be at a concentration near the midpoint of the curve. In many cases FGL uses a Laboratory Control Sample (LCS) as an ICV. In this case the LCS verifies both the calibration and sample preparation. FGL uses control charts for LCS's and acceptance ranges for many analytes have been statistically derived. Please see table 5-1 for acceptance limits. If calculated acceptance criteria are not listed, the general acceptance range is +/-25% of the true value for organics and +/-10% for inorganics.

8.3.10 Continuing Calibration Verifications (CCV)

The CCV is used to verify continuing calibration validity without having to completely restandardize the instrument. Refer to specific EPA methods or SOPs to determine whether this is required. The continuing calibration standard should be near the mid-point of the calibration curve. If calculated acceptance criteria are not listed, the general acceptance range is +/-25% of the true value for organics and +/-10% for inorganics.

8.3.11 Organic Mass Spectrometry Tuning Criteria

In addition to the above calibration criteria, GC/MS methods require mass spectrometer tuning. The acceptance criteria for volatile organics using bromofluorobenzene (BFB) are listed in table 8-1. The acceptance criteria for semivolatile organics using decafluorotriphenylphosphene (DFTPP) are listed in table 8-2.

Table 8-1

GC/MS Volatile Organic and Air Toxics Key Ion Abundance Tuning Criteria, using BFB

<u>Mass</u>	Ion Abundance Criteria
50	15 to 40% of mass 95
75	30 to 60% of mass 95
95	base peak, 100% relative abundance
96	5 to 9% of mass 95
173	less than 2% of mass 174
174	greater than 50% of mass 95
175	5 to 9% of mass 174
176	greater than 95% but less than 101% of mass 174
177	5 to 9% of mass 176

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Table 8-2

GC/MS Semivolatile Organic Key Ion Abundance Tuning Criteria, using DFTPP

<u>Mass</u>	<u> Ion Abundance Criteria</u>
51	30 to 60% of mass 198
68	less than 2% of mass 69
70	less than 2% of mass 69
127	40 to 60% of mass 198
197	less than 1% of mass 198
198	base peak, 100% relative abundance
199	5 to 9% of mass 198
275	10 to 30% of mass 198
365	greater than 1% of mass 198
441	present but less than mass 443
442	greater than 40% of mass 198
443	17 to 23% of mass 442

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Calibration Procedures and Frequency

8.3.12 Calibration Frequency

8.3.12.1 Inorganics

For trace metals analyses calibrations are performed initially for each analytical run. After initial calibration verification (ICV) recalibration or continuing calibration checks are used every ten analyses. For wet chem analyses, a new curve is generated every six months. To verify daily calibration of wet chem methods a continuing calibration verifications (CCV) is used. Exceptions may exist for method specific SOPs.

8.3.12.2 Organics

GC and GC/MS methods are calibrated initially for all analytes. For each additional day of operation, a calibration check standard is analyzed and evaluated. If the calibration is deemed valid the analysis may be performed. Otherwise, the system is recalibrated. Exceptions may exist for method specific SOPs.

8.3.12.3 Radiochemistry

All methods utilize recalibration or continuing calibration verifications every ten samples. EPA check samples are performed quarterly. DOE check sample are performed on a semiannual basis. The results must be acceptable to maintain certification. Exceptions may exist for certain method specific SOP's

8.3.12.3.1 Gas Proportional Counters

Gas proportional counters are calibrated for each matrix and sample size. For most liquid samples an efficiency vs. mg chart for each instrument is fit to a polynomial equation, which is stored in the computer for automated calculation.

8.3.12.3.2 Liquid Scintillation

Liquid Scintillation analyzers are calibrate for each matrix and sample size. Standards are initially run in duplicate with calibration verifications performed on each analytical run.

8.3.12.3.3 Gamma Spectroscopy

Gamma spectroscopy analyses are calibrated for both energy and efficiency. The calibration range is from 20 keV to 1350 keV with a minimum of 7 strong peaks per calibration for each geometry and matrix.

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Analytical Procedures

9.1 Method Sources

FGL uses EPA or standard methods for most analyses. The following method manuals are used as references:

Drinking Water Methods -

- 1) "Methods for Chemical Analysis in Waters and Waste," (MCAWW) EPA-600/4-79-020
- 2) "Standard Methods for the Analysis of Water and Wastewater," 17th Edition, 1990.
- 3) "Methods for the Determination of Organic Compounds in Drinking Water," EPA Method Book, EPA-600/4-88-039, December 1988.
- 4) "Methods for the Determination of Organic Compounds in Drinking Water-Supplement I," EPA Method Book, EPA-600/4-90-020, July 1990.
- 5) "Methods for the Determination of Organic Compounds in Drinking Water-Supplement II," EPA Method Book, EPA-600/4- 90-020, July 1990.
- 6) "Eastern Environmental Radiation Facility Radiochemistry Procedures Manual," EPA Method Book, EPA 520/5-84-006, August 1984.
- 7) "Environmental Measurements Laboratory Procedures," HASL-300, 27th Edition, February 1992.

Wastewater Methods -

- 1) "Methods for Chemical Analysis in Waters and Waste," (MCAWW) EPA-600/4-79-020
- 2) "Standard Methods for the Analysis of Water and Wastewater," 17th Edition, 1990.
- 3) "Methods for Organic Chemical Analysis of Municipal and Industrial Wastewater," EPA Method Book, EPA 600/4-82-057, July 1982.
- 4) "Eastern Environmental Radiation Facility Radiochemistry Procedures Manual," EPA Method Book, EPA 520/5-84-006, August 1984.
- 5) "Environmental Measurements Laboratory Procedures," HASL-300, 27th Edition, February 1992.

Solid Waste / Hazardous Waste Methods -

- 1) "Methods for Evaluating Solid Waste," EPA Method Book, SW- 846, revision 3, and proposed revisions.
- 2) "Eastern Environmental Radiation Facility Radiochemistry Procedures Manual," EPA Method Book, EPA 520/5-84-006, August 1984.
- 3) "Environmental Measurements Laboratory Procedures," HASL-300, 27th Edition, February 1992.

Ambient Air / Air Pollution Methods -

- 1) "Compendium of Methods for the Determination of Toxic Organic Compounds in Ambient Air," EPA Method Book, EPA-600/4-, June 1988.
- 2) "Technical Assistance Document for Sampling and Analysis of Toxic Organic Compounds in Ambient Air," EMSL, EPA Method Book, EPA-600/4-83-027, 1983

9.2 Standard Operating Procedures (SOPs)

FGL has specific standard operating procedures (SOPs) for each analytical test we perform. The specific format and use for these procedures is given in SOP S0QA010.XXX.

9.3 Specific Methods Used for Analysis

The analytical methods performed at FGL fall into three general categories: drinking waters, wastewaters, solid waste / hazardous waste and ambient air / air pollution methods. These methods are listed in Tables 9-1 - 9-4.

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TABLE 9-1 Specific Analytical Drinking Water Methods

Parameter	Method	Description
Organic Chemicals		
Trihalomethanes	EPA 501.2	GC/ECD, micro extraction
Volatile Organics	EPA 502.2	GC/PID/Hall, purge & trap
Dibromochloropropane (DBCP)	EPA 504	GC/ECD, micro extraction
and Ethylene dibromide (EDB)		,
Chlorinated Pesticides	EPA 505	GC/ECD, micro extraction
Nitrogen/phosphorus Pesticides	EPA 507	GC/NPD, liquid-liquid
Chlorothalonil	EPA 508	GC/ECD, liquid-liquid
PCB's as Decachlorobiphenyl	EPA 508A	GC/ECD, liquid-liquid
Trihalomethane Form. Potential	EPA 510	GC/ECD, micro extraction
Herbicides	EPA 515.1	GC/ECD, liquid-liquid
Volatile Organics	EPA 524.2	GC/MS, purge & trap
Diethylhexylphthalate	EPA 525	GC/MS, SPE
Carbamates	EPA 531	HPLC, post column derivatization
Glyphosate (Roundup)	EPA 547	HPLC, post column derivatization
Endothall	EPA 548	HPLC, post column derivatization
Paraquat and Diquat	EPA 549	HPLC, post column derivatization
Polynuclear Aromatic Hydrocarbons	EPA 550.1	HPLC, post column derivatization
Haloacetic Acids	EPA 552.1	HPLC, post column derivatization, SPE
General Inorganic Analyses		
Acidity	EPA 305.1	Titration
Aggressive Index	N/A	Calculation
Alkalinity (CaCO3)	EPA 310.1	Titration
Bicarbonate (HCO3)	EPA 310.1	Titration
Biochemical Oxygen Demand (BOD5)	EPA 405.1	ISE
Bromide (Br)	EPA 300.0	IC
Carbon Dioxide (CO2)	SM 4500CO2	Titration
Carbonate (CO3)	EPA 310.1	Titration
Chemical Oxygen Demand (COD)	EPA 410.2	Colorimetric
Chloride (Cl)	EPA 300.0	IC
Chlorine Residual (Cl2)	EPA 330.2	Titration
Chlorine Residual (Cl2)	EPA 330.5	Colorimetric
Chlorine Demand	SM 409A	Colorimetric
Color	EPA 110.3	Observation
Cyanide, Total (CN)	EPA 335.2	Colorimetric
Electrical Conductivity (EC)	EPA 120.1	Conductivity Bridge
Fluoride (F)	EPA 340.2	ISE
Hardness, total (as CaCO3)	EPA 130.2	Titration
Hardness, total (as CaCO3)	SM 2340B	Calculation
Hydroxide (OH)	EPA 310.1	Titration
Langelier Index (corrosivity)	SM 2330B	Calculation
MBAS	EPA 425.1	Colorimetric

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TABLE 9-1 Specific Analytical Drinking Water Methods

Parameter	Method	Description
General Inorganic Analyses continued	i	
Nitrogen		
Ammonia (NH3-N)	EPA 350.1	Colorimetric
Nitrate (NO3-N)	EPA 300.0	IC
Nitrite (NO2-N)	EPA 300.0	IC
Nitrate (NO3-N)	EPA 353.2	Colorimetric
Nitrite (NO2-N)	EPA 353.2	Colorimetric
Organic (TKN-NH3-N)	N/A	Calculation
Total (TKN+NO3-N+NO2-N)	N/A	Calculation
Total Kjeldahl	EPA 351.2	Colorimetric
Odor	EPA 140.1	Observation
Oil and Grease	EPA 413.1	Gravimetric
Oxygen, Dissolved (DO)	EPA 360.1	ISE
pH	EPA 150.1	ISE
Phenols	EPA 420.1	Colorimetric
Phosphorous	2-11-1-002	
Phosphate (PO4-P)	EPA 300.0	IC
Phosphate-dissolved (PO4-P)	EPA 300.0	IC
Total (P)	EPA 365.2	Colorimetric
Total-dissolved (P)	EPA 365.2	Colorimetric
Resistivity	N/A	Calculation
Sodium Percent	N/A	Calculation
Sodium Absorption Ratio (SAR)	EPA 200.7	ICP
Solids/Residue	EI A 200.7	101
Filterable (TDS)	EPA 160.1	Gravimetric
Non-filterable (TSS)	EPA 160.1	Gravimetric
Total	EPA 160.3	Gravimetric
Volatile	EPA 160.3	Gravimetric
Settleable	EPA 160.5	Gravimetric
	EPA 300.0	IC
Sulfate (SO4)	EFA 300.0	ic
Sulfide (H2S) Total	EDA 276 2	Mathylana Plua
Dissolved	EPA 376.2 EPA 376.2	Methylene Blue Methylene Blue
Sulfite (SO2)	EPA 370.2 EPA 377.1	•
,		Titrimetric
Tannin & Lignin	SM 5500B	Colorimetric
Titration - pH adjustment	N/A EDA 1901	Titration
Turbidity	EPA 180.1	Nephelometric
Trace Metals Analyses		
Aluminum (Al)	EPA 200.9	Furnace Atomic Absorption
	EPA 200.9	ICP/MS
Aluminum (Al)	EPA 200.8 EPA 200.9	
Antimony (Sb)		Furnace Atomic Absorption
Antimony (Sb)	EPA 200.8	ICP/MS
Arsenic (As)	EPA 200.9	Furnace Atomic Absorption
Arsenic (As)	EPA 200.8	ICP/MS
Barium (Ba)	EPA 200.7	ICP ACC
Barium (Ba)	EPA 200.8	ICP/MS

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TABLE 9-1 Specific Analytical Drinking Water Methods

Parameter	Method	Description
Trace Metals Analyses continued		
Beryllium (Be)	EPA 200.7	ICP
Beryllium (Be)	EPA 200.8	ICP/MS
Boron (B)	EPA 200.7	ICP
Boron (B)	EPA 200.8	ICP/MS
Cadmium (Cd)	EPA 200.9	Furnace Atomic Absorption
Cadmium (Cd)	EPA 200.8	ICP/MS
Calcium (Ca)	EPA 200.7	ICP
Chromium (Cr)	EPA 200.9	Furnace Atomic Absorption
Chromium (Cr)	EPA 200.8	ICP/MS
Chromium VI (Cr+6)	EPA 7196	Colorimetric
Cobalt (Co)	EPA 200.7	ICP
Cobalt (Co)	EPA 200.8	ICP/MS
Copper (Cu)	EPA 200.7	ICP
Copper (Cu)	EPA 200.8	ICP/MS
Iron (Fe)	EPA 200.7	ICP
Lead (Pb)	EPA 200.9	Furnace Atomic Absorption
Lead (Pb)	EPA 200.8	ICP/MS
Lithium (Li)	SM 3500LiB	Flame Atomic Absorption
Magnesium (Mg)	EPA 200.7	ICP
Manganese (Mn)	EPA 200.7	ICP
Manganese (Mn)	EPA 200.8	ICP/MS
Mercury (Hg)	EPA 245.1	Cold Vapor Atomic Absorption
Mercury (Hg)	EPA 245.2	Cold Vapor Atomic Absorption
Molybdenum (Mo)	EPA 200.7	ICP
Molybdenum (Mo)	EPA 200.8	ICP/MS
Nickel (Ni)	EPA 200.7	ICP
Nickel (Ni)	EPA 200.8	ICP/MS
Potassium (K)	EPA 200.7	ICP
Selenium (Še)	EPA 200.9	Furnace Atomic Absorption
Selenium (Se)	EPA 200.8	ICP/MS
Silica (SiO2)	EPA 200.7	ICP
Silver (Ag)	EPA 200.9	Furnace Atomic Absorption
Silver (Ag)	EPA 200.8	ICP/MS
Sodium (Na)	EPA 200.7	ICP
Thallium (Tl)	EPA 200.9	Furnace Atomic Absorption
Thallium (Tl)	EPA 200.8	ICP/MS
Tin (Sn)	EPA 200.9	Furnace Atomic Absorption
Vanadium (V)	EPA 200.7	ICP
Vanadium (V)	EPA 200.8	ICP/MS
Zinc (Zn)	EPA 200.7	ICP
Zinc (Zn)	EPA 200.7	ICP/MS

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TABLE 9-1 Specific Analytical Drinking Water Methods

Parameter	Method	Description
Radiochemical Analyses		
Gross Alpha	EPA 900.0	Proportional Counter
Gross Beta	EPA 900.0	Proportional Counter
Gross Alpha & Beta	EPA 900.0	Proportional Counter
Gamma Emmitters	EPA 901.1	HPĜe, gamma spectroscope
Total Radium*	EPA 900.1	Isolation, Proportional Counter
Radium 226	EPA 903.1	Radon bubbler, Lucas cell scintillation
Radium 228	EPA 904.0	Isolation, Proportional Counter
Uranium	EPA 908.0	Isolation, Proportional Counter
Tritium	EPA 906.0	Distillation, Liquid Scintillation
Radon	EPA 913.0	Liquid Scintillation
Bacteriological Analyses		
Total & Fecal Coliform	SM9221E	Fermentation, MPN, 10 tube
Total Coliform-Colilert	SM9221D	Presence-Absence
Standard Plate Count	SM9215B	Incubation, visual count

^{*} Can be reported as Radium 226 if less than 3 pCi/liter.

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TABLE 9-2 Specific Analytical Wastewater / Hazardous Waste Liquid Methods

Parameter Method Description	
Organic Chemical Analyses	
Sample Preparation EPA 3510 Liquid-Liquid	extraction
1 1	uid- Liquid Extraction
Sample Preparation EPA 3580 Solvent Dilution	
Purgeable Halocarbons EPA 601/8010 GC/PID/Hall,	
EDB and DBCP EPA 8011 GC/ECD, micro	
, , , , , , , , , , , , , , , , , ,	
71 0	
, , , , , , , , , , , , , , , , , , , ,	e & trap
Purgeable Non-Holographical Valetile Organics - EPA 8015M - CC/FID micro	a autuantian
Non-Halogenated Volatile Organics EPA 8015M GC/FID, micro	o extraction
Extractable FRA (02/9020 GG/FID TOWN	P. 4
Aromatic Volatile Organics EPA 602/8020 GC/FID, purg	
Phenols EPA 604/8040 GC/ECD, liqu	
Chlorinated Pesticides & PCB's EPA 608/8080 GC/ECD, liqu	
Polynuclear Aromatic Hydrocarbons EPA 610/8310 HPLC/UV, liq	
Organophosphorus Pesticides EPA 614/8140 GC/FPD, liqui	
Chlorinated Herbicides EPA 615/8150 GC/ECD, liqu	
Volatile Organics EPA 624/8240 GC/MS, purge	
Semi-volatile Organics EPA 625/8270 GC/MS, liquid	
Carbamates EPA 632 HPLC/UV, liq	
Total Organic Carbon (TOC) EPA415.1/9060 IR, combustio	
Total Organic Halogens (TOX) EPA 9020 Coulometric,	
Total Recov. Pet. Hydrocarbons EPA 418.1 IR, liquid-liqu	iid
General Inorganic Analyses	
Acidity EPA 305.1 Titration	
Aggressive Index N/A Calculation	
Alkalinity (CaCO3) EPA 310.1 Titration	
Bicarbonate (HCO3) EPA 310.1 Colorimetric	
Biochemical Oxygen Demand (BOD5) EPA 405.1 ISE	
Bromide (Br) EPA 300.0 IC	
Carbonate (CO3) EPA 310.1 Titration	
Carbon Dioxide (CO2) SM 4500CO2 Titration	
Chemical Oxygen Demand (COD) EPA 410.2 Colorimetric	
Chloride (Cl) EPA 300.0 IC	
Chlorine Residual (Cl2) EPA 330.2 Titration	
Chlorine Residual (Cl2) EPA 330.5 Colorimetric	
Chlorine Demand SM 409A Titration	
Color EPA 110.3 Visual	
Cyanide, Total (CN) EPA 335.2 Colorimetric	
Electrical Conductivity (EC) EPA 120.1 Conductivity	Bridge
Fluoride (F) EPA 340.2 ISE	
Hardness, total (as CaCO3) EPA 130.2 Titration	
Hydroxide (OH) EPA 310.1 Titration	
Langelier Index (corrosivity) SM 2330B Calculation	

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TABLE 9-2 Specific Analytical Wastewater / Hazardous Waste Liquid Methods

1		•
Parameter	Method	Description
C 11 ' A 1	1	
General Inorganic Analyses continue	ea	
Nitrogen	ED 4 250 1	Calanimatria
Ammonia (NH3-N)	EPA 350.1	Colorimetric
Nitrate (NO3-N)	EPA 300.0	IC
Nitrite (NO2-N)	EPA 300.0	IC
Organic (TKN-NH3-N)	Calculation	
Total (TKN+NO3-N+NO2-N)	Calculation	~
Total Kjeldahl	EPA 351.2	Colorimetric
Odor	EPA 140.1	Observation
Oil and Grease	EPA 413.1	Gravimetric
Oxygen, Dissolved (DO)	EPA 360.1	ISE
pH	EPA 150.1	ISE
Phenols	EPA 420.1	Colorimetric
Phosphorous		
Phosphate (PO4-P)	EPA 300.0	IC
Phosphate-dissolved (PO4-P)	EPA 300.0	IC
Total (P)	EPA 365.2	Colorimetric
Total-dissolved (P)	EPA 365.2	Colorimetric
Resistivity	N/A	Calculation
Sodium Percent	N/A	Calculation
Sodium Absorption Ratio (SAR)	EPA 200.7	ICP
Solids/Residue		
Filterable (TDS)	EPA 160.1	Gravimetric
Non-filterable (TSS)	EPA 160.2	Gravimetric
Total	EPA 160.3	Gravimetric
Volatile	EPA 160.4	Gravimetric
Settleable	EPA 160.5	Gravimetric
Sulfate (SO4)	EPA 300.0	IC
Sulfide (H2S)		
Total	EPA 376.2	Methylene Blue
Dissolved	EPA 376.2	Methylene Blue
Sulfite (SO2)	EPA 377.1	Titrimetric
Tannin & Lignin	SM 513	Colorimetric
Titration - pH adjustment	N/A	Titration
Turbidity	EPA 180.1	Nephelometric
•		•
Trace Metals Analyses		
Sample Preparation	EPA 3015	Digestion
Aluminum (Al)	EPA 200.9	Furnace Atomic Absorption
Aluminum (Al)	EPA 200.8	ICP/MS
Antimony (Sb)	EPA 200.9	Furnace Atomic Absorption
Antimony (Sb)	EPA 200.8	ICP/MS
Arsenic (As)	EPA 200.9	Furnace Atomic Absorption
Arsenic (As)	EPA 200.8	ICP/MS
Barium (Ba)	EPA 200.7	ICP
Barium (Ba)	EPA 200.8	ICP/MS
Beryllium (Be)	EPA 200.7	ICP
Beryllium (Be)	EPA 200.8	ICP/MS
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TABLE 9-2 Specific Analytical Wastewater / Hazardous Waste Liquid Methods

Parameter	Method	Description
Trace Metals Analyses continued		
Boron (B)	EPA 200.7	ICP
Boron (B)	EPA 200.8	ICP/MS
Cadmium (Cd)	EPA 200.9	Furnace Atomic Absorption
Cadmium (Cd)	EPA 200.8	ICP/MS
Calcium (Ca)	EPA 200.7	ICP
Chromium (Cr)	EPA 200.9	Furnace Atomic Absorption
Chromium (Cr)	EPA 200.8	ICP/MS
Chromium VI (Cr+6)	EPA 7196	Colorimetric
Cobalt (Co)	EPA 200.7	ICP
Cobalt (Co)	EPA 200.8	ICP/MS
Copper (Cu)	EPA 200.7	ICP
Copper (Cu)	EPA 200.8	ICP/MS
Gold (Au)	EPA 231.1	Flame Atomic Absorption
Iron (Fe)	EPA 200.7	ICP
Lead (Pb)	EPA 200.9	Furnace Atomic Absorption
Lead (Pb)	EPA 200.8	ICP/MS
Lithium (Li)	SM 3500LiB	Flame Atomic Absorption
Magnesium (Mg)	EPA 200.7	ICP
Manganese (Mn)	EPA 200.7	ICP
Manganese (Mn)	EPA 200.8	ICP/MS
Mercury (Hg)	EPA 245.1	Cold Vapor Atomic Absorption
Mercury (Hg)	EPA 245.2	Cold Vapor Atomic Absorption
Molybdenum (Mo)	EPA 200.7	ICP
Molybdenum (Mo)	EPA 200.8	ICP/MS
Nickel (Ni)	EPA 200.7	ICP
Nickel (Ni)	EPA 200.8	ICP/MS
Potassium (K)	EPA 200.7	ICP
Selenium (Še)	EPA 200.9	Furnace Atomic Absorption
Selenium (Se)	EPA 200.8	ICP/MS
Silica (SiO2)	EPA 200.7	ICP
Silver (Ag)	EPA 200.9	Furnace Atomic Absorption
Silver (Ag)	EPA 200.8	ICP/MS
Sodium (Na)	EPA 200.7	ICP
Strontium (Sr)	EPA 200.7	ICP
Thallium (Tl)	EPA 200.9	Furnace Atomic Absorption
Thallium (Tl)	EPA 200.8	ICP/MS
Tin (Sn)	EPA 200.9	Furnace Atomic Absorption
Titanium (Ti)	EPA 200.7	ICP
Uranium (Ù)	EPA 200.7	ICP
Vanadium (V)	EPA 200.7	ICP
Zinc (Zn)	EPA 200.7	ICP
Zinc (Zn)	EPA 200.7	ICP/MS

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TABLE 9-2 Specific Analytical Wastewater / Hazardous Waste Liquid Methods

Parameter	Method	Description
Radio Chemical Analyses		
Gross Alpha	EPA 900.0	Proportional counter
Gross Beta	EPA 900.0	Proportional counter
Gross Alpha & Beta	EPA 900.0	Proportional counter
Gamma Emmitters	EPA 901.1	HPGe, gamma spectroscope
Total Radium*	EPA 900.1	Isolation, Proportional Counter
Radium 226	EPA 903.1	Radon bubbler, Lucas cell scintillation
Radium 228	EPA 904.0	Isolation, Proportional Counter
Uranium	EPA 908.0	Isolation, proportional counter
Tritium	EPA 906.0	Distillation, liquid scintillation
Radon	EPA 913.0	Liquid scintillation
Bacteriological Analyses		
Total Coliform	SM9221E	Fermentation, MPN, 15 tube
Total & Fecal Coliform	SM9221E	Fermentation, MPN, 15 tube
Standard Plate Count	SM9215B	Incubation, visual count

^{*} Can be reported as Radium 226 if less than 3 pCi/liter.

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TABLE 9-3 Specific Analytical Solid Waste / Hazardous Waste Methods

Parameter	Method	Description
Organic Chemical Analyses		
Sample Preparation - STLC	Title 22	Extraction
Sample Preparation - EP TOX	EPA 1310	Extraction
Sample Preparation - TCLP	EPA 1311	Extraction
Sample Preparation	EPA 3510	Liquid-Liquid extraction
Sample Preparation	EPA 3520	Continous Liquid- Liquid Extraction
Sample Preparation	EPA 3540	Soxhlet Extraction
Sample Preparation	EPA 3550	Sonication Extraction
Sample Preparation	EPA 3580	Solvent Dilution
Purgeable Halocarbons	EPA 8010	GC/PID/Hall, purge & trap
EDB and DBCP	EPA 8011	GC/ECD, micro extraction
Non-Halogenated Volatile Organics	EPA 8015	GC/FID, purge & trap
Total Petroleum Hydrocarbons	EPA 8015M	GC/FID, purge & trap
Purgeable	ED 4 001534	CC/FID
Total Petroleum Hydrocarbons Extractable	EPA 8015M	GC/FID, micro extraction
Aromatic Volatile Organics	EPA 8020	GC/FID, purge & trap
Phenols	EPA 8040	GC/CED, soxhlet or sonication
Chlorinated Pesticides & PCB's	EPA 8080	GC/ECD, soxhlet or sonication
Polynuclear Aromatic Hydrocarbons	EPA 8100	GC/PID, soxhlet or sonication
Organophosphorus Pesticides	EPA 8141	GC/FPD, soxhlet or sonication
Chlorinated Phenoxy Herbicides	EPA 8150	GC/ECD, soxhlet or sonication
Volatile Organics	EPA 8240	GC/MS, purge & trap
Semi-volatile Organics	EPA 8270	GC/MS, soxhlet or sonication
Polynuclear Aromatic Hydrocarbons	EPA 8310	HPLC/UV, liquid- liquid
Carbamates	EPA 632	HPLC/UV, liquid- liquid
Total Organic Carbon (TOC)	EPA 9060	IR, combustion
Total Organic Halogens (TOX)	EPA 9020	Coulometric, Pyrolysis
Total Recov. Pet. Hydrocarbons	EPA 418.1	IR, liquid-liquid
General Inorganic Analyses		
Chloride (Cl)	EPA 9056	IC
Electrical Conductivity	EPA 120.1	Conductivity bridge
Cyanide, total (CN)	EPA 335.2	Distillation- Colorimetric
Fluoride (F)	EPA 340.1	Distillation-ISE
Moisture	ASA/UL	Gravimetric
Nitrogen	77. 460. 4	
Ammonia (NH3-N)	EPA 350.1	Colorimetric
Nitrate (NO3-N)	EPA 9056	IC
Nitrite (NO2-N)	EPA 9056	IC
Organic (TKN-NH3-N)	N/A	Calculation
Total (TKN+NO3-N+NO2-N)	N/A	Calculation
Total Kjeldahl	EPA 351.1	Colorimetric
Oil and grease	EPA 413.1M	Gravimetric
Soxhlet	EFA 413.1WI	GI AVIIIICU IC

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TABLE 9-3 Specific Analytical Solid Waste / Hazardous Waste Methods

Parameter	Method	Description
General Inorganic Analyses continued		
pH	EPA 9045	ISE
Phosphorous	TD + 0056	10
Phosphate (PO4)	EPA 9056	IC
Total (P)	See Trace Metals	
Sulfate (SO4)	EPA 9056 EPA 376.2	IC Colorimetric
Sulfide (H2S)	EPA 5/0.2	Colorimetric
Hazardous Waste Characterization A	nalyses	
Corrosivity (pH)		
Aqueous sample	EPA 9040	ISE
Nonaqueous sample	EPA 9045	ISE
Ignitability	ED 4 1010	T1 1 1 1
Aqueous (Flashpoint)	EPA 1010	Flashpoint
Nonaqueous (Flammability)	EPA 1020	Flashpoint
Reactivity	SW-846 Ch 8	Observations
Reaction with water		
Reaction with dilute acid		
Reaction with dilute base		
Reaction with oxidizing agent		
Reaction with reducing agent Generation	SW-846 Ch 7	Screens
Sulfide	5 W-040 CH 7	Sciens
Cyanide		
Cyamuc		
Trace Metals	ED . 4050	D.
Sample Preparation - TTLC	EPA 3050	Digestion
Sample Preparation - STLC	Title 22	Extraction
Sample Preparation - EP TOX	EPA 1310	Extraction
Sample Preparation - TCLP	EPA 1311	Extraction
Aluminum (Al)	EPA 6010 EPA 6020	ICP ICP/MS
Aluminum (Al) Antimony (Sb)	EPA 7041	Furnace Atomic Absorption
Antimony (Sb)	EPA 6020	ICP/MS
Arsenic (As)	EPA 7060	Furnace Atomic Absorption
Arsenic (As)	EPA 6020	ICP/MS
Barium (Ba)	EPA 6010	ICP
Barium (Ba)	EPA 6020	ICP/MS
Beryllium (Be)	EPA 6010	ICP
Beryllium (Be)	EPA 6020	ICP/MS
Boron (B)	EPA 6010	ICP
Boron (B)	EPA 6020	ICP/MS
Cadmium (Cd)	EPA 6010	ICP
Cadmium (Cd)	EPA 6020	ICP/MS
Calcium (Ca)	EPA 6010	ICP
Chromium (Cr)	EPA 6010	ICP
Chromium (Cr)	EPA 6010	ICP/MS
Chromium VI (Cr+6)	EPA 7196	Colorimetric

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TABLE 9-3 Specific Analytical Solid Waste / Hazardous Waste Methods

Trace Metals Analyses continued Cobalt (Co)	Parameter	Method	Description
Cobalt (Co) EPA 6010 ICP Cobalt (Co) EPA 6020 ICP/MS Copper (Cu) EPA 6020 ICP/MS Copper (Cu) EPA 6020 ICP/MS Gold (Au) EPA 231.1 Flame Atomic Absorption Iron (Fe) EPA 6010 ICP Lead (Pb) EPA 7420 Flame Atomic Absorption Lead (Pb) EPA 6020 ICP/MS Lithium (Li) EPA 7430 Flame Atomic Absorption Magnesium (Mg) EPA 6010 ICP Manganese (Mn) EPA 6010 ICP Manganese (Mn) EPA 6010 ICP Manganese (Mn) EPA 6010 ICP Molybdenum (Mo) EPA 6010 ICP Nickel (Ni) EPA 6010 ICP Potassium (K) EPA 6010 ICP Potassium (K) EPA 6010 ICP Selenium (Se) EPA 7741 Furnace Atomic Absorption Selenium (Se) EPA 6020 ICP/MS Silver (Ag) EPA 6020 ICP/MS Silver (Ag) EPA 6020 ICP/MS Silver (Ag) EPA 6010 ICP Silver (Ag) EPA 6010 ICP Strontium (Sr) EPA 6010 ICP Thallium (Tl) EPA 6010 ICP Thallium (Tl) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP Uranium (U) EPA 6010 ICP Vanadium (V) EPA 6010 ICP	Trace Metals Analyses continued		
Copper (Cu) EPA 6010 ICP Copper (Cu) EPA 6020 ICP/MS Gold (Au) EPA 231.1 Flame Atomic Absorption Iron (Fe) EPA 6010 ICP Lead (Pb) EPA 7420 Flame Atomic Absorption Lead (Pb) EPA 6020 ICP/MS Lithium (Li) EPA 6020 ICP/MS Lithium (Li) EPA 6010 ICP Manganese (Mn) EPA 6010 ICP Manganese (Mn) EPA 6010 ICP Manganese (Mn) EPA 6020 ICP/MS Mercury (Hg) EPA 7470 Cold Vapor Atomic Absorption Molybdenum (Mo) EPA 6010 ICP Molybdenum (Mo) EPA 6010 ICP Molybdenum (Mo) EPA 6010 ICP Nickel (Ni) EPA 6010 ICP Nickel (Ni) EPA 6010 ICP Selenium (Se) EPA 6010 ICP Selenium (Se) EPA 6010 ICP Selenium (Se) EPA 6010 ICP Silver (Ag) EPA 6010 ICP Silver (Ag) EPA 6010 ICP Strontium (Sr) EPA 6010 ICP Strontium (Tl) EPA 6010 ICP Titanium (Tl) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP Vanadium (V) EPA 6010 ICP	•	EPA 6010	ICP
Copper (Cu) EPA 6010 ICP Copper (Cu) EPA 6020 ICP/MS Gold (Au) EPA 231.1 Flame Atomic Absorption Iron (Fe) EPA 6010 ICP Lead (Pb) EPA 7420 Flame Atomic Absorption Lead (Pb) EPA 6020 ICP/MS Lithium (Li) EPA 6020 ICP/MS Lithium (Li) EPA 6010 ICP Manganese (Mn) EPA 6010 ICP Manganese (Mn) EPA 6010 ICP Manganese (Mn) EPA 6020 ICP/MS Mercury (Hg) EPA 7470 Cold Vapor Atomic Absorption Molybdenum (Mo) EPA 6010 ICP Molybdenum (Mo) EPA 6010 ICP Molybdenum (Mo) EPA 6010 ICP Nickel (Ni) EPA 6010 ICP Nickel (Ni) EPA 6010 ICP Selenium (Se) EPA 6010 ICP Selenium (Se) EPA 6010 ICP Selenium (Se) EPA 6010 ICP Silver (Ag) EPA 6010 ICP Silver (Ag) EPA 6010 ICP Strontium (Sr) EPA 6010 ICP Strontium (Tl) EPA 6010 ICP Titanium (Tl) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP Vanadium (V) EPA 6010 ICP		EPA 6020	ICP/MS
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Magnesium (Mg)EPA 6010ICPManganese (Mn)EPA 6010ICPManganese (Mn)EPA 6020ICP/MSMercury (Hg)EPA 7470Cold Vapor Atomic AbsorptionMolybdenum (Mo)EPA 6010ICPMolybdenum (Mo)EPA 6020ICP/MSNickel (Ni)EPA 6010ICPNickel (Ni)EPA 6020ICP/MSPhosphorous (P)EPA 6010ICPSelenium (Se)EPA 6010ICPSelenium (Se)EPA 6020ICP/MSSilver (Ag)EPA 6010ICPSilver (Ag)EPA 6010ICPSodium (Na)EPA 6010ICPStrontium (Sr)EPA 6010ICPThallium (Tl)EPA 7841Furnace Atomic AbsorptionThallium (Tl)EPA 6010ICPTin (Sn)EPA 6010ICPTitanium (Ti)EPA 6010ICPUranium (U)EPA 6010ICPVanadium (V)EPA 6010ICPZinc (Zn)EPA 6010ICP		EPA 7430	Flame Atomic Absorption
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Manganese (Mn) Mercury (Hg) EPA 6020 EPA 7470 Cold Vapor Atomic Absorption Molybdenum (Mo) EPA 6010 ICP Molybdenum (Mo) EPA 6020 ICP/MS Nickel (Ni) EPA 6010 ICP Nickel (Ni) EPA 6020 ICP/MS Phosphorous (P) Potassium (K) EPA 6010 EPA 6010 ICP Selenium (Se) EPA 7741 Furnace Atomic Absorption Selenium (Se) EPA 6020 ICP/MS Silver (Ag) EPA 6010 ICP Silver (Ag) EPA 6010 ICP Strontium (Sr) EPA 6010 ICP Thallium (Tl) EPA 6010 ICP Thallium (Tl) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP		EPA 6010	ICP
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Molybdenum (Mo) Nickel (Ni) Nickel (Ni) EPA 6010 EPA 6020 ICP/MS ICP/MS Phosphorous (P) Potassium (K) Selenium (Se) Selenium (Se) Silver (Ag) Silver (Ag) Sodium (Na) Sodium (Na) Strontium (Sr) Thallium (Tl) Thallium (Tl) Thallium (Tl) EPA 6010 ICP Titanium (Tl) EPA 6010 ICP		EPA 7470	Cold Vapor Atomic Absorption
Molybdenum (Mo) Nickel (Ni) Nickel (Ni) Phosphorous (P) Potassium (K) Selenium (Se) Selenium (Se) Silver (Ag) Silver (Ag) Sodium (Na) Strontium (Sr) Thallium (Tl) Thallium (Tl) Titanium (Ti) Uranium (U) Vanadium (V) Zinc (Zn) EPA 6010 EPA 6010 EPA 6010 ICP ICP/MS ICP ICP ICP ICP ICP ICP ICP IC	• • • •	EPA 6010	
Nickel (Ni) EPA 6010 ICP Nickel (Ni) EPA 6020 ICP/MS Phosphorous (P) EPA 6010 ICP Potassium (K) EPA 6010 ICP Selenium (Se) EPA 7741 Furnace Atomic Absorption Selenium (Se) EPA 6020 ICP/MS Silver (Ag) EPA 6010 ICP Silver (Ag) EPA 6020 ICP/MS Sodium (Na) EPA 6010 ICP Strontium (Sr) EPA 6010 ICP Thallium (Tl) EPA 7841 Furnace Atomic Absorption Thallium (Tl) EPA 6020 ICP/MS Tin (Sn) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP Uranium (U) EPA 6010 ICP Vanadium (V) EPA 6010 ICP Zinc (Zn) EPA 6010 ICP		EPA 6020	ICP/MS
Phosphorous (P) Potassium (K) EPA 6010 EPA 6010 ICP Selenium (Se) EPA 7741 Furnace Atomic Absorption Selenium (Se) EPA 6020 ICP/MS Silver (Ag) EPA 6010 ICP Silver (Ag) EPA 6010 ICP Strontium (Sr) EPA 6010 ICP Strontium (Tl) EPA 6010 ICP Thallium (Tl) EPA 6020 ICP/MS ICP Titanium (Tl) EPA 6010 ICP Titanium (Tl) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP Uranium (U) EPA 6010 ICP Uranium (U) EPA 6010 ICP Cinc (Zn) EPA 6010 ICP		EPA 6010	ICP
Phosphorous (P) Potassium (K) EPA 6010 EPA 6010 ICP Selenium (Se) EPA 7741 Furnace Atomic Absorption Selenium (Se) EPA 6020 ICP/MS Silver (Ag) EPA 6010 ICP Silver (Ag) EPA 6010 ICP Strontium (Sr) EPA 6010 ICP Strontium (Tl) EPA 6010 ICP Thallium (Tl) EPA 6020 ICP/MS ICP Titanium (Tl) EPA 6010 ICP Titanium (Tl) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP Uranium (U) EPA 6010 ICP Uranium (U) EPA 6010 ICP CICP Cinc (Zn) EPA 6010 ICP		EPA 6020	ICP/MS
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Selenium (Se) Selenium (Se) EPA 7741 Furnace Atomic Absorption EPA 6020 ICP/MS Silver (Ag) Silver (Ag) Sodium (Na) EPA 6010 EPA 6010 ICP Strontium (Sr) Thallium (Tl) EPA 7841 EPA 6020 Teph Surnace Atomic Absorption EPA 6020 Thallium (Tl) EPA 6020 Teph Surnace Atomic Absorption Thallium (Tl) EPA 6020 Ticp/MS Tin (Sn) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP Uranium (U) EPA 6010 ICP Uranium (V) EPA 6010 ICP Ticp	EPA 6010	ICP	
Selenium (Se) EPA 6020 ICP/MS Silver (Ag) EPA 6010 ICP Silver (Ag) EPA 6020 ICP/MS Sodium (Na) EPA 6010 ICP Strontium (Sr) EPA 6010 ICP Thallium (Tl) EPA 7841 Furnace Atomic Absorption Thallium (Tl) EPA 6020 ICP/MS Tin (Sn) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP Uranium (U) EPA 6010 ICP Vanadium (V) EPA 6010 ICP Zinc (Zn) EPA 6010 ICP		EPA 7741	Furnace Atomic Absorption
Silver (Ag) Silver (Ag) Silver (Ag) EPA 6020 EPA 6020 ICP/MS Sodium (Na) EPA 6010 ICP Strontium (Sr) EPA 6010 EPA 7841 Furnace Atomic Absorption Thallium (Tl) EPA 6020 ICP/MS Tin (Sn) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP Uranium (U) EPA 6010 ICP Uranium (V) EPA 6010 ICP Uranium (V) EPA 6010 ICP ICP ICP Vanadium (V) EPA 6010 ICP ICP ICP ICP ICP ICP ICP IC	Selenium (Se)	EPA 6020	ICP/MS
Silver (Ag) Sodium (Na) EPA 6020 ICP/MS Sodium (Na) EPA 6010 ICP Strontium (Sr) EPA 6010 ICP Thallium (Tl) EPA 7841 Furnace Atomic Absorption Thallium (Tl) EPA 6020 ICP/MS Tin (Sn) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP Uranium (U) EPA 6010 ICP Vanadium (V) EPA 6010 ICP Zinc (Zn) EPA 6010 ICP		EPA 6010	ICP
Sodium (Na) Strontium (Sr) Thallium (Tl) EPA 6010 EPA 7841 Furnace Atomic Absorption Thallium (Tl) EPA 6020 ICP/MS Tin (Sn) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP Uranium (U) EPA 6010 ICP Uranium (V) EPA 6010 ICP Uranium (V) EPA 6010 ICP Uranium (V) EPA 6010 ICP ICP ICP ICP ICP ICP ICP IC	Silver (Ag)	EPA 6020	ICP/MS
Thallium (Tl) EPA 7841 Furnace Atomic Absorption EPA 6020 ICP/MS Tin (Sn) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP Uranium (U) EPA 6010 ICP Uranium (V) EPA 6010 ICP ICP ICP Vanadium (V) EPA 6010 ICP ICP ICP ICP ICP ICP ICP IC		EPA 6010	ICP
Thallium (Tl) EPA 7841 Furnace Atomic Absorption EPA 6020 ICP/MS Tin (Sn) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP Uranium (U) EPA 6010 ICP Uranium (V) EPA 6010 ICP ICP ICP Vanadium (V) EPA 6010 ICP ICP ICP ICP ICP ICP ICP IC		EPA 6010	ICP
Tin (Sn) EPA 6010 ICP Titanium (Ti) EPA 6010 ICP Uranium (U) EPA 6010 ICP Vanadium (V) EPA 6010 ICP Zinc (Zn) EPA 6010 ICP		EPA 7841	Furnace Atomic Absorption
Titanium (Ti) EPA 6010 ICP Uranium (U) EPA 6010 ICP Vanadium (V) EPA 6010 ICP Zinc (Zn) EPA 6010 ICP	Thallium (Tl)	EPA 6020	ICP/MS
Uranium (U) EPA 6010 ICP Vanadium (V) EPA 6010 ICP Zinc (Zn) EPA 6010 ICP	Tin (Sn)	EPA 6010	ICP
Vanadium (V) EPA 6010 ICP Zinc (Zn) EPA 6010 ICP	Titanium (Ti)	EPA 6010	ICP
Zinc (Zn) EPA 6010 ICP	Uranium (U)	EPA 6010	ICP
7inc (7n) FPA 6020 ICP/MS	Zinc (Zn)		
Zine (Zii)	Zinc (Zn)	EPA 6020	ICP/MS

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TABLE 9-3 Specific Analytical Solid Waste / Hazardous Waste Methods

Parameter	Method	Description
Radio Chemical Analyses		
Gross Alpha	EPA 900.0	Proportional counter
Gross Beta	EPA 900.0	Proportional counter
Gross Alpha & Beta	EPA 900.0	Proportional counter
Gamma Emmitters	EPA 901.1	HPĜe, gamma spectroscope
Total Radium*	EPA 900.1	Isolation, Proportional Counter
Radium 226	EPA 903.1	Radon bubbler, Lucas cell scintillation
Radium 228	EPA 904.0	Isolation, Proportional Counter
Uranium	EPA 908.0	Isolation, proportional counter
Tritium	EPA 906.0	Distillation, liquid scintillation
Radon	EPA 913.0	Liquid scintillation

^{*} Can be reported as Radium 226 if less than 3 pCi/liter.

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TABLE 9-4 Specific Analytical Ambient Air / Air Pollution Methods

Parameter	Method	Description
Volatile Organics	TO-14	GC/MS
BTEX/TPH	TO-14 (mod)	GC/MS
Total Non-methane hydrocarbons	TO-14 (mod)	GC/MS
Sulfur compounds	TO-14 (mod)	GC/MS
Light Hydrocarbons (C1-C6)	TO-3 (mod)	GC/FID
Methane	3810/5020	GC/FID
Fixed Gases	ASTM 1945	GC/TCD
Hydrogen Sulfide	-	Gold Film

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Data Reduction, Validation, and Reporting

The process of transforming raw analytical data into a finished report involves steps which are generally grouped into the categories of data reduction, data validation, and reporting. It involves mathematical modeling of the standard calibration curves, statistical analysis of the acquired data, calculations to account for preparation steps and dilutions, verification of adherence to quality assurance procedures, and the generation of hardcopy output.

10.1 Data Reduction

At FGL Environmental the analyst has the primary responsibility for reducing raw data. This process consists primarily of converting raw data into final reportable values by comparing individual sample results against those obtained for calibration purposes then accounting for any dilutions or concentration.

For each method, all raw data results are recorded on method specific forms or in a standardized output from each of the various instruments. Details on procedures for data reduction may be found in the laboratory SOP for each method.

10.2 Data Validation

Upon completion of each analytical run, the analyst enters or transfers the data to LIMS. The analytical raw data and LIMS generated QC summary sheets are validated by the laboratory supervisor or a backup peer analyst. They verify that all quality control parameters meet the data quality objectives and also review the analytical data for calculation errors and inconsistencies. When data quality objectives are not met the deficiency is investigated and the samples are rerun or an explanation is provided.

10.3 Data Review and Reporting Policy

The raw data review includes all documentation associated with the samples, including chromatograms, instrument run logs, digestion logs, and other instrument printouts. Upon approval by the analyst or supervisor, the analytical results for the run are transferred to a results database for compiling with other data for that sample. When all results for a sample have been entered, an on screen report is generated for review and validation by the supervisor. Upon approval by the supervisor sample results are then released for final hardcopy reporting, which is forwarded to the client. Quality assurance reports are also generated at this step. The final reports are reviewed before signing by both the department supervisor and laboratory director. Quality assurance reports are reviewed by the quality assurance director or officer. Invoices are also reviewed and initialed by the project manager or department supervisor.

Data review steps include the following:

- 1) All data packages are reviewed by a second analyst or the supervisor. The QC batch report and analytical run sheets (if applicable) must be initialed by the reviewer.
- 2) All supervisors must review the data released for reporting.
- 3) Analysis reports are printed and again reviewed by the supervisor and lab director and signed by each upon approval.
- 4) Quality Control reports are printed and are reviewed and signed by the quality assurance director or officer.
- 5) Invoices are reviewed and initialed by the project manager or department supervisor.

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Data Reduction, Validation and Reporting

10.4 Data Storage

FGL Environmental maintains report files and the supporting raw data for the current and previous year on the premises. Reports and raw data are maintained for a total of ten years in the secured data storage facility.

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Internal Quality Control Checks

An internal quality control program requires a set of routine internal procedures for assuring that the data generated from a measurement system meets prescribed criteria for data quality. An effective internal QC program must be capable of measuring and controlling the quality of the data, in terms of precision, accuracy, and completeness (see sections 5 and 14 for these details).

This section identifies QC protocols associated with analytical procedures. Tables 11-1 - 11-4 are general outlines of quality control parameters monitored for each procedure. Included are general quality control measures as well as specific quality control checks which provide continual control and assessment of data quality. Figures 11-1 through 11-3 are examples of FGL Control Charts. FGL uses continuing calibration verifications (CCV) and initial calibration verification (ICV) for instrument quality control. The laboratory control sample (LCS) is used for sample preparation quality control. The CCV standard only verifies continuing calibration. The ICV standard is used to independently verify the calibration and may take the place of CCV when used on a continuing basis during analysis. The LCS may take the place of both ICV and CCV when prepared independently and used on a continuing basis during analysis.

11.1 Quality Control Parameters

11.1.1 Initial Demonstration of Capability

Before analyzing samples, the laboratory must prove proficiency in the method by preparing a data package for certification. The laboratory normally provides the following information:

- 1) calibration data
- 2) calibration verification from an independent source
- 3) method detection limit data
- 4) detection limit verification data
- 5) accuracy and precision data

These must all be acceptable under the method QC criteria or, when requirements are not specified, reasonably meet good laboratory practices and Department of Health Services requirements.

11.1.2 Analysis Quality Controls

11.1.2.1 Instrument Blank

The instrument or calibration blank is used to calibrate the instrument. This blank contains the same reagents used in the standards and samples. However, the blank is prepared under controlled conditions and is not processed like all samples.

11.1.2.2 Initial Calibration Verification (ICV)

ICV is used as an independent verification of the calibration. EPA protocol recommends analysis of ICV for each analytical calibration. ICV samples are manufacturer or laboratory prepared from independent suitable reference standards. The ICV usually contains the analytes of interest at a concentration in the mid-calibration range. Method specific acceptance criteria are used when possible. When method specific criteria are unavailable, the recoveries are control charted to obtain acceptance limits.

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Internal Quality Control Checks

11.1.2.3 Continuing Calibration Verification (CCV)

CCV is used to verify continuing calibration. Many EPA methods require analysis of CCV's on a per batch or per day basis. CCV samples are manufacturer or laboratory prepared from suitable reference standards. This standard may or may not be independent of the calibration stock standards. The CCV usually contains the analytes of interest at a concentration in the mid-calibration range. Method specific acceptance criteria are used when possible. When method specific criteria are unavailable the recoveries are control charted to obtain acceptance limits.

11.1.2.4 Internal Standards (IS)

An IS is a synthetic compound not occuring in an environmental sample but has chemical behavior similar to that of the target analytes. EPA protocol requires IS for specific methods on a per sample basis (including QC samples). The IS serves as a check on the analysis and corrects for instrumental drift or matrix effects. Method specific acceptance criteria are used when possible. When method specific criteria are unavailable the recoveries are control charted to obtain acceptance limits.

11.1.3 Method Quality Controls

11.1.3.1 Method Blank

The method blank is used to ensure that any positive results were not because of reagent or labware contamination. Before analyzing any samples, the analyst must demonstrate through the analysis of a method blank, that all glassware and reagents are free of contaminants. Each time a set of samples is extracted, a method blank must be processed to check for laboratory contamination. The blank samples should be carried through all stages of the sample preparation and analysis. Lack of contamination is demonstrated if all target analytes with the exception of common laboratory reagents are below their DLRs.

11.1.3.2 Field Blank

The field blank is used to ensure that any positive results were not because of contamination occurring during sampling. The field blank samples should be carried through all stages of the sample preparation and analysis. Lack of contamination is demonstrated if all target analytes with the exception of common laboratory reagents are below their DLRs.

11.1.3.3 Travel Blank

The travel blank is used to ensure that any positive results were not because of contamination occurring during shipping and handling of the samples. The travel blank samples should be carried through all stages of the sample preparation and analysis. Lack of contamination is demonstrated if all target analytes with the exception of common laboratory reagents are below their DLRs.

11.1.3.4 Laboratory Control Sample (LCS)

The LCS is used to verify overall accuracy of the method. EPA protocol requires analysis of an LCS for each analytical batch when appropriate. The LCS consists of either a control matrix spiked with analytes representative of the target analytes or a certified reference material. Whenever possible, the LCS contains the analyte of interest at a concentration in the mid-calibration range. This standard may or may not be independent of the calibration stock standards. Initially method specific acceptance criteria are used. Eventually, or when method specific criteria are unavailable, the recoveries are control charted to obtain acceptance limits.

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Internal Quality Control Checks

11.1.3.5 Surrogate Spikes

Surrogate spikes serve as a check on the extraction process where extraction is a necessary part of the analytical procedure. When surrogate recovery is within limits it indicates that the extraction was complete. A surrogate is a compound not expected to occur in an environmental sample but has chemical behavior similar to that of the target analytes. EPA protocol requires surrogate spikes for specific methods on a per sample basis (including QC samples). Initially method specific acceptance criteria are used. Eventually, or when method specific criteria are unavailable, the recoveries are control charted to obtain acceptance limits.

11.1.3.6 Matrix Spike/Matrix Spike Duplicates (MS/MSD)

The MS/MSD is used to verify matrix specific precision and accuracy. EPA protocol normally requires analysis of MS/MSD samples for each analytical batch or matrix type. The MS/MSD spikes are manufacturer or laboratory prepared from suitable reference standards. This standard may or may not be independent of the calibration stock standards. The matrix spike recovery and relative percent difference (RPD) acceptance criteria are shown in Section 5. When matrix spike results fall outside limits published in the respective methods, The LCS is used to verify method control. If spike recoveries are outside normal limits due to matrix problems, the data should be reported noting matrix interference. The spike recovery and RPD acceptance limits are test specific and are control charted.

11.1.3.7 Duplicates

Duplicates are used to verify matrix specific precision. EPA protocol normally requires analysis of duplicate samples for each analytical batch or matrix type. The relative percent difference (RPD) calculated from duplicate analyses provide an assessment of precision. The RPD acceptance limits are test specific and are control charted.

11.1.4 Radiochemical Specific Quality Controls

11.1.4.1 Efficiency vs. Dissolved Solids Chart

Dissolved solids (TDS) mask or decrease the radiation picked in the proportional counters. For each instrument an efficiency vs. solids chart must be generated as part of the initial demonstration of capability. In addition, whenever an instrument is maintained or repaired (i.e. a counting wire replaced) a new efficiency vs. TDS chart must be generated. Samples containing solids such that the efficiency of a counter could drop below ten percent must be reprepared using a smaller aliquot so that the solids give acceptable counting efficiency. Whenever possible an electrical conductivity measurement is used to estimate TDS and sample aliquots.

11.1.4.2 Background

Background samples are run daily, prior to sample analysis. However, monthly average may be used for calculation purposes. If a run background is used for sample calculation the background must be within 2 standard deviations of the monthly average to be acceptable.

11.1.4.3 Minimum Detectable Activity (MDA)

MDA's are calculated every six months. This data is used to determine if a sample is not detectable. EPA guidelines for sensitivity are followed for each isotope. MDA's can be calculated on per sample basis. Background samples are run daily, prior to sample analysis. However, monthly average may be used for calculation purposes.

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Internal Quality Control

Table 11-1 Quality Controls for Drinking Water Methods

	INST	METHOD								
TEST	BLANK	BLANK	CCV	ICV	LCS	BS/BSD	MS/MSD	DUP	SURR	I.S.
Semivolatile Organics										
EPA 501.2	Batch	Batch		Batch	Batch					
EPA 504	Batch	Batch		Batch	Batch					
EPA 505	Batch	Batch		Batch	Batch					
EPA 507	Batch	Batch		Batch	Batch		Sample			
EPA 508	Batch	Batch		Batch	Batch		Sample			
EPA 508A	Batch	Batch		Batch	Batch		•			
EPA 510	Batch	Batch		Batch	Batch					
EPA 515.1	Batch	Batch		Batch	Batch		Sample			
EPA 525	Batch	Batch		Batch	Batch		Sample	Sample		
EPA 531	Batch	Batch		Batch	Batch		Sample	•		
EPA 547	Batch	Batch		Batch	Batch		•			
EPA 548	Batch	Batch		Batch	Batch		Sample			
EPA 549	Batch	Batch		Batch	Batch		•			
EPA 550.1	Batch	Batch		Batch	Batch		Sample			
EPA 552	Batch	Batch		Batch	Batch		Sample			
TOC	Batch		Batch		Batch		•			
TOX	Batch		Batch		Batch					
Volatile Organics										
EPA 502.2	Batch		Batch		Batch		Sample	Sample		
EPA 524.2	Batch		Batch		Batch		Sample	Sample		
Inorganic Chemicals			D 4 1			Datak				
Alkalinity			Batch		D-4-L	Batch				
Ammonia	Batch		Batch	Datab	Batch	Datak				
BOD	Batch			Batch		Batch				
Carbon Dioxide	D '1			Datak	Datah	Batch				
COD	Daily			Batch	Batch					
COD, % Transmittance		Datab		Batch	Batch		Batch			
Cl Res., colorimetric	Daily	Batch		Batch			Batch			
Cl Res., titrimetric				Batch			Datell			

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Table 11-1 Quality Controls for Drinking Water Methods

	INST	METHOD								• •
TEST	BLANK	BLANK	CCV	ICV	LCS	BS/BSD	MS/MSD	DUP	SURR	I.S.
Inorganic Chemical con	itinued									
Cyanide, Free	Batch		Batch		Batch					
Cyanide, Total	Batch	Batch	Batch		Batch	Batch				
E.C.			Batch			Batch				
Fluoride (Distillation)		Batch	Batch		Batch	Batch				
Fluoride	Batch		Batch		Batch					
General Physical										
Ignitability			Batch			Batch				
Ion Chromatography	Batch		Batch		Batch					
MBAS Extraction	Batch			Batch	Batch					
MBAS Screen							Batch			
Moisture, percent							Batch			
Oxygen, Dissolved							Batch			
Nitrate - Technicon	Batch		Batch		Batch					
Nitrite - Technicon	Batch		Batch		Batch					
Oil & Grease, Pet	Batch			Batch	Batch					
Oil & Grease, Soxhlet	Batch			Batch	Batch					
Oil & Grease	Batch			Batch	Batch					
р Н					Batch			Batch		
pH, Adjustment							Batch			
Phenols	Batch	Batch		Batch	Batch					
Phosphorous, Total	Batch	Batch		Batch	Batch					
Reactivity, Generation							Batch			
Reactivity							Batch			
Solids, Fixed	Batch			Batch		Batch				
Solids, Settleable										
Solids, Total	Batch			Batch		Batch				
Solids, T. Dissolved	Batch			Batch		Batch				
Solids, T. Suspended	Batch			Batch		Batch				
Solids, Volatile	Batch				Batch					
Solids, V. Suspended	Batch				Batch					
Sulfide, Dissolved							Batch			

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Internal Quality Control

Table 11-1 Quality Controls for Drinking Water Methods

TEST	INST BLANK	METHOD BLANK	CCV	ICV	LCS	BS/BSD	MS/MSD	DUP	SURR	I.S.
TEST	DLANK	BLANK	CCV	icv	LCS	BS/BSD	MS/MSD	БСТ	Som	1.01
Inorganic Chemical cor	ntinued									
Sulfide, Total								Batch		
Sulfite								Batch		
TKN	Batch				Batch		Batch			
Tannin & Lignin	Batch				Batch		Batch			
Trace Metals										
Cr VI	Batch		Batch		Batch		Batch			
FAA	Batch		Batch				Batch			
GFAA	Batch		Batch				Batch			
ICP	Batch		Batch				Batch			
ICP/MS	Batch		Batch				Batch			Sample
Hg - CVAA	Batch		Batch		Batch		Batch			
Radiochemistry										
Gross A & B		Batch						Batch		
Gamma		Batch						Batch		
Nuclide Screen		Batch				Batch				
Radium 226		Batch					Batch			
Radium 228		Batch				5	Batch			
Radium 226 & 228		Batch				Batch		D-4-1		
Radon		Batch					D 4 a b	Batch		
Strontium 90		Batch				D-4-1	Batch			
Tritium		Batch				Batch	D . I			
Uranium		Batch					Batch			

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Table 11-2 Quality Controls for Wastewater / Hazardous Waste Liquid Methods

	INST	METHOD								
TEST	BLANK	BLANK	CCV	ICV	LCS	BS/BSD	MS/MSD	DUP	SURR	1.S.
Semivolatile Organics										
EPA 3510				Batch			Batch	Batch		
EPA 3520				Batch			Batch	Batch		
EPA 3540				Batch			Batch	Batch		
EPA 3550				Batch			Batch	Batch		
EPA 3580				Batch			Batch			Batch
TCLP				Batch						Batch
EPA 8015	Batch			Batch						
EPA 8015M (Purgeable)	Batch			Batch						
EPA 8015M (Extract)	Batch			Batch						
EPA 604/8040	Batch			Batch						Sample
EPA 608/8080	Batch			Batch					Sample	
EPA 610/8310	Batch			Batch						
EPA 614/8140	Batch			Batch						Sample
EPA 615/8150	Batch			Batch						
EPA 625/8270	Batch			Batch						Sample
EPA 632	Batch			Batch						
TOC	Batch			Batch			Batch			
TOX	Batch			Batch			Batch			
TPH by IR	Batch	Batch			Batch	Batch				
Volatile Organics										
EPA 601/8010	Batch			Batch			Batch		Sample	Sample
EPA 602/8020	Batch			Batch			Batch		Sample	-
EPA 624/8240	Batch			Batch			Batch		Sample	Sample
EFA 024/0240	Daten			Duten					·	_
Inorganic Chemicals							D 4 1			
Alkalinity			Batch				Batch			
Ammonia	Batch			Batch			Batch	Datah		
BOD	Batch				Batch			Batch		Batch
Carbon Dioxide						D . 1	D . 4 . b			Daten
COD		Daily				Batch	Batch			

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Internal Quality Control

Table 11-2 Quality Controls for Wastewater / Hazardous Waste Liquid Methods

TEST	INST BLANK	METHOD BLANK	ccv	ICV	LCS	BS/BSD	MS/MSD	DUP	SURR	I.S.
Inorganic Chemicals con	ntinued									
COD, % Transmittance					Batch	Batch				
CI Res., colorimetric	Batch			Batch				Batch		
CI Res., titrimetric					Batch				Batch	
Cyanide, Free	Batch			Batch			Batch			
Cyanide, Total	Batch	Batch	Batch		Batch	Batch				
E.C.			Batch				Batch			
Fluoride (Distillation)			Batch	Batch			Batch	Batch		
Fluoride	Batch			Batch			Batch			
Gen. Physical										Batch
lgnitability					Batch					Batch
Ion Chromatography	Batch			Batch			Batch			
MBAS Extraction	Batch				Batch	Batch				
MBAS Screen									Batch	
Moisture, percent										Batch
Oxygen, Dissolved									Batch	
Nitrate - Technicon	Batch			Batch			Batch			
Nitrite - Technicon	Batch			Batch			Batch			
Oil & Grease, Pet	Batch				Batch	Batch				
Oil & Grease, Sox	Batch				Batch	Batch				
Oil & Grease	Batch				Batch	Batch				
рН						Batch				Batch
pH, Adjustment						Б.,				Batch
Phenols	Batch	Batch			Batch	Batch				
Phosphorus, Total	Batch	Batch			Batch	Batch			B	
Reactivity, Generation									Batch	D. A. I
Reactivity	D 4-1				Datah			Dotah		Batch
Solids, Fixed	Batch				Batch			Batch		
Solids, Settleable	Dotal				Dotah			Batch		
Solids, Total	Batch				Batch Batch			Batch		
Solids, T. Dissolved	Batch							Batch		
Solids, T. Suspended	Batch				Batch			Daten		

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Table 11-2 Quality Controls for Wastewater / Hazardous Waste Liquid Methods

TEST	INST BLANK	METHOD BLANK	CCV	ICV	LCS	BS/BSD	MS/MSD	DUP	SURR	I.S.
Inorganic Chemicals co Solids, Volatile Solids, V. Suspended Sulfide, Diss Sulfide, Total Sulfite TKN Tannin & Lignin	ntinued Batch Batch Batch Batch				Batch Batch	Batch Batch	Batch Batch			Batch Batch Batch
Trace Metals 3015 STLC TCLP Cr VI FAA GFAA ICP ICP/MS Hg - CVAA	Batch Batch Batch Batch Batch Batch	Batch	Batch	Batch Batch Batch Batch Batch Batch Batch	Batch	Batch	Batch Batch Batch	Batch		Batch Sample
Radiochemistry Gross A & B Gamma Nuclide Screen Radium 226 Radium 228 Radon Radium 226 & 228 Strontium 90 Tritium Uranium	Batch	Batch Batch Batch Batch Batch Batch Batch	Batch			Batch Batch Batch Batch Batch	Batch Batch Batch	Batch Batch		

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Internal Quality Control

Table 11-3 Quality Controls for Solid Waste / Hazardous Waste Methods

TEST	INST BLANK	METHOD BLANK	CCV	ICV	LCS	BS/BSD	MS/MSD	DUP	SURR	I.S.
Semivolatile Organics										
EPA 3510				Batch			Batch	Batch		
EPA 3510 EPA 3520				Batch			Batch	Batch		
EPA 3540				Batch			Batch	Batch		
EPA 3550				Batch			Batch	Batch		
EPA 3580				Batch			Batch	Daten		Batch
TCLP				Batch			Daten			Batch
EPA 8015	Batch			Batch						Daten
EPA 8015M (Diesel)	Batch			Batch						
EPA 8015M (Gas)	Batch			Batch						Sample
				Batch						Sample
EPA 8040	Batch Batch			Batch						Sample
EPA 8080				Batch						Sample
EPA 8140	Batch			Batch						Sample
EPA 8150	Batch									
EPA 8310	Batch			Batch						
EPA 632	Batch			Batch				•		
EPA 8270	Batch			Batch						
TOC	Batch			Batch			Batch			
TOX	Batch			Batch			Batch			
TPH by IR	Batch	Batch		Batch		Batch				
Volatile Organics										
TCLP				Batch						Batch
EPA 601/8010	Batch			Batch			Batch		Sample	Sample
EPA 602/8020	Batch			Batch			Batch		Sample	-
EPA 624/8240	Batch			Batch			Batch		Sample	Sample
EPA 8260	Batch			Batch			Batch		Sample	Sample
DI AL UZUU										•

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Table 11-3 Quality Controls for Solid Waste / Hazardous Waste Methods

TEST	INST BLANK	METHOD BLANK	ccv	ICV	LCS	BS/BSD	MS/MSD	DUP	SURR	1.S.
Solids Inorganic Chemi Ammonia Corrosivity (pH)	icals Batch				Batch Batch	Batch			Batch	
Cyanide, Total	Batch	Batch	Batch		Batch		Batch			
E.C. Fluoride (Distillation)			Batch Batch	Batch			Batch Batch	Batch		
Ignitability			Dute		Batch					Batch
Ion Chromatography	Batch				Batch	Batch				Batch
Moisture, percent Nitrate - Technicon	Batch				Batch	Batch				
Nitrite - Technicon	Batch				Batch	Batch				
Oil & Grease, Sox	Batch				Batch	Batch				
Oil & Grease	Batch				Batch	Batch				Batch
рН					Batch	D . I				Daten
Phenols	Batch	Batch			Batch	Batch				
Phosphorus (See Trace	metals - ICP)							Batch	
Reactivity, Generation									Daten	Batch
Reactivity									Batch	Daten
Solids, Percent									Daten	Batch
Sulfide, Total					Batch	Batch	Batch			
TKN	Batch				Daten	Daten	Daten			
Trace Metals			5.1			Batch	Batch			
3050			Batch	D.4.b		Batch	Daten			Batch
STLC				Batch			Batch	Batch		2,41011
TCLP				Batch	Batch		Batch	Daten		
Cr VI	Batch	Batch		Batch	Daten		Daten			
FAA	Batch			Batch						
GFAA	Batch			Batch						
ICP	Batch			Batch						Sample
ICP/MS	Batch			Batch						•
Hg - CVAA	Batch			Daten						

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Table 11-3 Quality Controls for Solid Waste / Hazardous Waste Methods

TEST	INST BLANK	METHOD BLANK	CCV	ICV	LCS	BS/BSD	MS/MSD	DUP	SURR	1.0
1631	DLANK	DLAINK	CCV	icv	LCS	D3/D3D	MIS/MISD	DOF	SURK	1.S.
Radiochemistry										
Gross A & B		Batch			Batch					
Gamma		Batch				Batch				
Nuclide Screen		Batch			Batch					
Radium 226		Batch			Batch					
Radium 228		Batch			Batch					
Radium 226 & 228		Batch			Batch					
Radon		Batch				Batch				
Strontium 90		Batch			Batch					
Tritium		Batch			Batch					
Uranium		Batch			Batch					

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Table 11-4 Quality Controls for Ambient Air / Air Pollution Methods

TEST	INST BLANK	METHOD BLANK	ccv	ICV	LCS	BS/BSD	MS/MSD	DUP	SURR	1.S.
TO-14 TO-3 (modified) Fixed Gases	Batch		Batch Batch Batch		Batch Batch Batch			Batch Batch Batch	Sample	Sample
Total Pet. Hydrocarbon Hydrogen Sulfide	s Batch Batch		Batch		Batch			Batch Batch	Sample	Sample

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Internal Quality Control Checks

Figure 11-1 FGL Control Chart for LCS

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QALCS-LBW

FGL ENVIRONMENTAL TREND CONTROL CHART

300.0 Anions for CL

09/23/93

BATCH	LOW	% RECOVERD (n= 30) AVG 107	HIGH
ID	98		115
09/17/93:B 09/17/93:A 09/16/93:C 09/16/93:B 09/16/93:A 09/15/93:B 09/15/93:A 09/13/93:B 09/13/93:A 09/10/93:D 09/10/93:C 09/10/93:C 09/10/93:B 09/10/93:A 09/08/93:C 09/08/93:C 09/08/93:A 09/03/93:C 09/03/93:A 09/02/93:B 09/02/93:B 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C	Д		۵

△/■ => Matrix Spike / Dup

Reporting Limits: 98-115 MAV % on 9/23/93 by: 6 / 10/18/93 BA

QALCS-LBW

FGL ENVIRONMENTAL TREND CONTROL CHART

300.0 Anions for CL

09/23/93

BATCH	DATE	Theo.	MS	MSD	AR	See	
ID	COMPLETED	Conc.	%	%	%	Notes	
09/17/93:B 09/17/93:A 09/16/93:C 09/16/93:B 09/15/93:B 09/15/93:A 09/13/93:B 09/13/93:A 09/10/93:D 09/10/93:C 09/10/93:C 09/10/93:A 09/10/93:A 09/08/93:C 09/08/93:B 09/08/93:A 09/03/93:A 09/02/93:B 09/02/93:A 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C 09/01/93:C	09/17/93 09/17/93 09/16/93 09/16/93 09/16/93 09/15/93 09/15/93 09/13/93 09/13/93 09/10/93 09/10/93 09/10/93 09/10/93 09/10/93 09/08/93 09/08/93 09/08/93 09/03/93 09/03/93 09/02/93 09/02/93 09/01/93 09/01/93 09/01/93 09/01/93 08/30/93 08/27/93 08/27/93 08/27/93 08/27/93	213 213 213 213 213 213 213 213 213 213	105 101 110 109 105 108 105 104 106 110 108 107 106 108 105 105 105 105 107 107 95.2 112 109 121 109 108 106 110	75-125 75-125	1		

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Internal Quality Control Checks

Figure 11-2 FGL Control Chart for MS/MSD and RPD

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QAMS-WW

FGL ENVIRONMENTAL TREND CONTROL CHART

300.0 Anions for CL

09/23/93

BATCH ID	LOW 54	% RECOVERD (n= 60) AVG 102	HIGH 150	% DIFFERENCE AVG 2.4	(n= 30) MAV 7.9
09/17/93:B		۵			
09/16/93:C		۵			
09/16/93:B		<u> </u>	<u> </u>		
09/15/93:B		i	[_		
09/13/93:B		△ ■	-		
09/10/93:D					
09/10/93:C		△ •			
09/10/93:B		O=			
09/08/93:C		0		+	
09/08/93:B		• ۵		→	
09/03/93:C		•		↔	
09/02/93:B		Δ .			
09/02/93:A	-	Δ			-
09/01/93:C	,	■ △			
09/01/93:B		Δ •		.	
08/30/93:B		_ _ •		↔	
08/30/93:A		■ b		↔	
08/27/93:F		Ţ.		→	
08/27/93:C		•			
08/26/93:E		i - △		↔	
08/26/93:C		a		↔	
08/24/93:E		i △ =		 ↔	
08/24/93:D		•△		•	
08/24/93:C		•			
08/24/93:B		■ □		•	
08/20/93:F		Δ■		•	
08/18/93:E		Δ=		↔	
08/18/93:D		•		↔	
08/18/93:C		- 4		↔	
08/18/93:B		ф•		+	

△/■ => Matrix Spike / Dup

↔ => % Difference

Reporting Limits: S4-150 7.9 on 9 /33/93 by: C / 10/18/93 \$A

QAMS-WW

FGL ENVIRONMENTAL TREND CONTROL CHART

300.0 Anions for CL

09/23/93

BATCH ID	DATE COMPLETED	Theo. Conc.	MS %	AR %	DIFF. %	MAV %	See Notes	
09/17/93:B	09/17/93	100	98.9	101	75-125	0.6	20.0	10
09/16/93:C	09/16/93	40.0	96.9	99.3	75-125	0.5	20.0	*
09/16/93:B	09/16/93	100	154	151	75-125	1.2	20.0	*
09/15/93:B	09/15/93	145	180	180	75-125	0.1	20.0	*
09/13/93:B	09/13/93	100	92.0	120	75-125	13.4	20.0	^
09/10/93:D	09/10/93	100 200	110 98.9	116 103	75-125 75-125	2.4	20.0 20.0	
09/10/93:C 09/10/93:B	09/10/93 09/10/93	100	106	103	75-125 75 - 125	1.6 1.6	20.0	
09/10/93:B 09/08/93:C	09/10/93	100	110	118	75-125 75-125	2.4	20.0	10
09/08/93:B	09/08/93	200	124	113	75-125 75-125	2.4	20.0	10
09/03/93:C	09/03/93	100	95.7	95.8	75-125	0.1	20.0	
09/02/93:B	09/02/93	100	95.0	102	75-125	4.5	20.0	
09/02/93:A	09/02/93	100	68.6	46.0	75-125	14.1	20.0	10
09/01/93:C	09/01/93	200	85.7	82.0	75-125	3.3	20.0	13
09/01/93:B	09/01/93	200	84.5	89.7	75-125	2.8	20.0	
08/30/93:B	08/30/93	100	98.5	103	75-125	1.3	20.0	
08/30/93:A	08/30/93	100	103	100	75-125	1.7	20.0	
08/27/93:F	08/27/93	4000	45.7	38.0	75-125	2.0	20.0	10
08/27/93:C	08/27/93	100	107	106	75-125	0.4	20.0	10
08/26/93:E	08/26/93	200	112	104	75-125	3.2	20.0	10
08/26/93:C	08/26/93	40.0	98.9	100	75-125	0.8	20.0	
08/24/93:E	08/24/93	50.0	109	111	75-125	1.3	20.0	
08/24/93:D	08/24/93	625	97.2	95.3	75-125	1.9	20.0	
08/24/93:C	08/24/93	250	106	106	75-125	0.6	20.0	13
08/24/93:B	08/24/93	250	109	107	75-125	1.2	20.0	
08/20/93:F	08/20/93	50.0	77.9	80.3	75-125	1.0	20.0	
08/18/93:E	08/18/93	250	83.8	86.4	75-125	1.5	20.0	13
08/18/93:D	08/18/93	100	99.5	99.7	75-125	0.1	20.0	10
08/18/93:C 08/18/93:B	08/18/ 9 3 08/18/93	100 40.0	101 102	94.7 103	75-125 75-125	3.0 0.5	20.0 20.0	10 13

		•	6 895 :
			1989 : 1985 -
			1989 : 1984 1985

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Performance and System Audits

The Quality Assurance Director or Officer is responsible for internal system audits, coordinating all external audits and performance evaluation (PE) samples. In addition, the QA Director or Officer is responsible for maintaining state and agency certification.

12.1 System Audits

System audits are performed both by external agencies, and by the laboratory Quality Assurance officers. These audits check the overall laboratory "system", from login to delivery of the finished reports. The purpose of the audit is to verify and document compliance with the regulatory methods, our Standard Operating Procedures (SOPs) and client specific needs.

12.1.1 Internal System Audit Program

Internal system audits are conducted on a monthly basis. Several analytical methods are selected each month for vertical slice audits. Compliance with all the required QC is evaluated and indicated on the QA Audit form (document ID F2QA040.XXX for Santa Paula, figure 12-1, and F3QA040.XXX for Stockton). The findings from the audit are compiled in the monthly QA non-conformance summary report.

12.1.2 External System Audits

System audits are performed by outside government agencies such as the California Department of Health Services, Lawrence Livermore National Laboratory and the Army Corp of Engineers. Audits are also performed by private agencies such as Chemical Waste Management, Inc.

12.2 Performance Evaluation Samples

Performance evaluation audits are used to provide a direct evaluation of the ability of the analytical systems to generate data that is consistent with the laboratory's stated objectives for accuracy and precision. External PE samples are analyzed as part of the certification and approval process for various state and federal agencies, as well as for other organizations.

12.2.1 External Performance Evaluation Samples

Performance evaluation samples are analyzed for a number of outside agencies including:

- (1) USEPA semi-annual drinking water check samples (WS series)
- (2) USEPA semi-annual wastewater check samples (WP series)
- (3) USEPA annual wastewater check samples (DMR studies)
- (4) CA-DHS-ELAP semi-annual hazardouswaste check samples (HW studies)
- (5) EMSL, Las Vegas, radiochemistry check samples
- (6) DOE, Environmental Measurements Laboratory QA Program
- (7) DOE, Mixed Analyte Performance Evaluation Program (MAPEP)

12.3 Certifications, Accreditations and Agency Approvals

FGL Environmental participates in laboratory certification programs with California, and other states. A copy of the California Environmental Laboratory Accredititation Program (ELAP) certified analysis list may be found in Figures 12-2 and 12-3 for Santa Paula and Stockton Laboratories respectively. A copy of the Utah and Hawaii Department of Health certified analysis lists may be found in Figures 12-4 and 12-5 (respectively) for Santa Paula.

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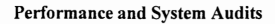


Figure 12-1 FGL QA Audit Form

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FGL - Santa Paula QA Audit Form

Date:		
Inspector:		
<u>Item</u>	In Compliance?	Corrective Action
Overall - Oven Temps Ref. Temps Balance Checks Container Checks QC SOPs		
Comments:		
Front Office Login Sample #s -	_	
LIMS Logbook File Tracking Completeness Chain of Custody Report Bill Training SOPs		
Comments:		
Bacteriology -		
MUG Log (colilert) MTF Log (D.W.) Series Log (W.W.) Plate Count Log Client Notifications Media Prep Log Autoclave Temps Water Suitability Corrective Actions Method Check		
	Sample	#:Date Started:
SOP Analyst Training		
Comments:		

<u>Item</u>	In Compliance?	Corrective Action
Inorganic Wet Chen	nistry -	
Instrument Maintenar Dionex DX-100 Dionex DX-300 P.E. Lambda 3 Turner Nepth. Fisher pH/ISE Orion E.C. Orion BOD Skalar BOD Corrective Actions	nce:	
Method Checks		
Method: Data Package Batch QC Report Reviewed Report Batch Sheet Inst. Printout SOP Control Charts MDL Study Standard Tracking Analyst Training	Sample #	Date Started:
Method:	Sample #	Date Started:

<u>Item</u>	In Compliance?	Corrective Action
Inorganic Trace Me	tals -	
Instrument Maintenar P.E. 5000 P.E. 5100 TJA Atomscan 16 TJA Trace Anal. Fisons PQ-II Leeman PS200 HP-4500 ICP-MS Corrective Actions	nce:	
Method:	Sample #	Date Started:
Organics - GC Lab - Instrument Maintena GC 2 GC 3 GC 4 GC 5 GC 7 GC 8 Hitachi HPLC HP HPLC		
IR Corrective Actions		
Comments:		

<u>Item</u>	In Compliance?	Corrective Action
Method Check		
Data Package Batch QC Report Reviewed Report Batch Sheet Inst. Printout SOP Control Charts MDL Study Standard Tracking Analyst Training		Date Started:
GC/MS Lab: Instrument Maintenar GC 9 GC/MS 1 GC/MS 2 GC/MS 3 GC/MS 4 GC/MS 5 MCI TOX Astro TOC Corrective Actions	nce:	
Method Check Method: Data Package Batch QC Report Reviewed Report Batch Sheet Inst. Printout SOP Control Charts MDL Study Standard Tracking Analyst Training Comments:	Sample #	Date Started:

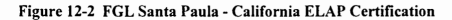
<u>Item</u>	In Compliance?	Corrective Action
Radiochemistry -		
Instrument Maintenant Alpha 1 Alpha 2 Alpha 5 Alpha 6 Alpha 7 Alpha/Beta 3 Alpha/Beta 4 Alpha/Beta 8 Liquid Scintillation Gamma Spec. Alpha Spec. Random Scint. Cntr. Corrective Actions Method Check		
Method: Data Package Batch QC Report Reviewed Report Batch Sheet Inst. Printout SOP Control Charts MDL Study Standard Tracking Analyst Training	Sample #	:Date Started:
Comments:		

<u>Item</u>	In Compliance?	Corrective Action
Agronomics Lab -		
Instrument Maintenar Technicon - nitrate Technicon - TKN LECO pH meter E.C. meter ARL 3410 Corrective Actions	nce:	
Method Check		
Data Package Batch QC Report Reviewed Report Batch Sheet Inst. Printout SOP	Sample #	Date Started:
Control Charts MDL Study Standard Tracking Analyst Training Comments:		

<u>Item</u>	In Compliance?	Corrective Action
Air Analysis Lab -		
Instrument Maintenar SRI GC/TCD SRI GC/FID HP 5890 GC/FID HP GC/MS #1 HP GC/MS #2 H ₂ S Analyzer Corrective Actions	nce:	
Method Check		
Method: Data Package Batch QC Report Reviewed Report Batch Sheet Inst. Printout SOP Control Charts MDL Study Standard Tracking Analyst Training Comments:	Sample #	Date Started:

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Performance and System Audits



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ENVIRONMENTAL LABORATORY ACCREDITATION/REGISTRATION List of Approved Fields of Testing and Analytes

FGL Environmental Laboratory 853 Corporation Street Santa Paula, CA TELEPHONE No: (805) 659-0910 CALIFORNIA COUNTY: Ventura

CERTIFICATE NUMBER: 1573 EXPIRATION DATE: 7/31/97

1	Microbiology of Drinking Water and Wastewater (
•						
1.1	Total Coliforms in Drinking Water by Multiple Tub	oe Ferment	tation Y			
1.2	Facal Californe/E Cali in Drinking Water by MTE		,			
1.3	Total Coliforms in Drinking Water by Membrane Fil	lter Techr	nics N			
1.4	Fecal Coliforms/E. Coli in Drinking Water by Memb	orane filt	ter TechnicsN			
1.5	Total Coliforms and E. Coli in Drinking Water by	MMO-MUG -	Y			
1.6	Total Coliforms in Drinking Water by Clark's Pres Fecal Coliforms/E. Coli in Drinking Water by Clar	sence/Abse	ence N			
1.7	Heterotrophic Plate Count	rk's Prese	ence/ Absence			
1.8 1.9	Total Coliforms in Wastewater by Multiple Tube Fo	ermentatio	nn Y			
1.10	Feral Coliforme in Wastewater by MTF		Y			
1.11	Total Coliforme in Wastewater by Membrane Filter	Technics				
1.12	Feral Coliforms in Wastewater by Membrane Filter	Technics	N			
1.13	Fecal Strentococci or Enterococci by Multiple Tul	he Technia	re			
1.14	Fecal Streptococci or Enterococci by Membrane Fi	lter Techi	nics N			
_						
2	Inorganic Chemistry and Physical Properties of	<u>Drinking</u>	Water excluding Toxic Chemical Elements			
	(07-15-91)					
2.1	Alkalinity Y	2.12	Sulfate Y			
2.2	CalciumY	2.13	Total Filterable Residue			
2.3	Chloride Y		and Conductivity			
2.4	Corrosivity Y	2.14	Iron (Colorimetric Methods Only) N			
2.5	Fluoride Y	2.15	Manganese (Colorimetric Methods Only) - N			
2.6	Hardness ····· Y	2.16	Phosphate, ortho Y			
2.7	Magnesium Y	2.17	Silica (Colorimetric Methods Only) N			
2.8	MBAS Y Nitrate Y	2.18	Cyanide Y			
2.9	Nitrite Y					
2.10 2.11	Sodium Y					
2.11	Journal Control of the Control of th					
3	Analysis of Toxic Chemical Elements in Drinking	Water (07-15-91)			
3.1	Arsenic Y	3.11	Silver Y			
3.2	Barium Y	3.12	Zinc Y			
3.3	Cadmium Y	3.13	Aluminum ····· Y			
3.4	Chromium, total Y	3.14	Asbestos N EPA Method 200.7 Y			
3.5 3.6	Iron Y	3.15 3.16	EPA Method 200.8 (Unregulated Elements			
3.7	LeadY	3.10	and Lead Only) Y			
3.8	Manganese Y	3.17	Antimony Y			
3.9	Mercury Y	3.18	Bervllium Y			
3.10	Selenium Y	3.19	Nickel Y			
		3.20	Thattium Y			
4	Organic Chemistry of Drinking Water (measureme	nt by GC/	MS combination) (07-15-91)			
4.1	EPA Method 501.3		u			
4.2	EPA Method 524.2		Υ			
4.3	EPA Method 525		· · · · · · · · · · · · · · · · · · ·			
4.4	EPA Method 513	• • • • • • • • •	N			
5	Organic Chemistry of Drinking Water (excluding	measurem	ments by GC/MS combination) (07-15-91)			
	EPA Method 501.1 N	- 4/	EPA Method 548 Y			
5.1	EPA Method 501.2 Y	5.16	EPA Method 549 Y			
5.2 5.3	EPA Method 502.1 N	5.17 5.18	EPA Method 550 N			
5.4	EPA Method 502.2 Y	5.19	EPA Method 550.1 N			
5.5	EPA Method 503.1 N	5.20	EPA Method 551 Y			
5.6	EPA Method 504 Y	5.21	EPA Method 552 Y			
5.7	EPA Method 505 Y					
5.8	EPA Method 506 N					
5.9	EPA Method 507 Y					
5.10	EPA Method 508 Y					
5.11	EPA Method 508AY EPA Method 510.1N					
5.12 5.13	EPA Method 510.1 N EPA Method 515.1 Y					
5.14	EPA Method 531.1 Y					
5.15	EPA Method 547 Y					

CERTIFICATE NUMBER: 1573 EXPIRATION DATE: 7/31/97

6	Radiochemistry (07-15-91)		
6.1	Gross Alpha and Beta Radiation Y	6.11	Gross Alpha by Co-precipitation
6.2	Total Radium Y	6.12	Radium 228
6.3	Radium 226 Y	6.13	Radioactive Iodine
6.4	UraniumY		Gross Alpha & Beta in Hazardous Wastes)
	uranium · · · · · · · · · · · · · · · · · · ·	6.14	Gross Alpha & Beta in mazardous wastes i
6.5	Radon 222 Y	6.15	Alpha Emitting Radium Isotopes
6.6	Radioactive Cesium N		in Haz. Wastes
6.7	Iodine 131 N	6.16	Radium 228 in Hazardous Wastes
6.8	Radioactive Strontium Y		
6.9	Tritium Y		
6.10	Gamma and Photon Emitters Y		
0.10	damina and Photon Emitters		
7	Shellfish Sanitation ()		
7.1	Shellfish meat Microbiology		
7.2	Paralytic Shellfish Poison		
7.3	Domoic Acid		
	bollote Acid		
8	Aquatic Toxicity Bioassays ()		
8.1	Hazardous Waste Aquatic Toxicity Bioassay (Title	22, CCR,	66261.24(a)(6))
8.2	Wastewater Testing According to Kopperdahl (1976)	using Fr	eshwater fish 🛉
8.3	Wastewater Testing According to EPA/600/4-85/013	usina Fre	eshwater and/or Marine Organisms
8.4	Uastewater Testing by FPA Method 1000 0		
8.5	Wastewater Testing by EPA Method 1002.0		
	Wastewater Testing by EPA Method 1002.0		
8.6	Wastewater lesting by EPA Method 1003.0		
8.7	Wastewater Testing by EPA Method 1006		
8.8	Wastewater Testing by EPA Method 1007		
8.9	Wastewater Testing by EPA Method 1009		
8.10	Wastewater Testing According to Anderson, et. al.	(1990)	using Giant Kelp (Macrocystis pyrifera) I
8.11	Wastewater Testing According to Anderson, et. al.		
8.12	Wastewater Testing According to Dinnel and Stober		
0.12	(Strongylocentrotus purpuratus)	(1907)	using purple sea ordina
	(Strongylocentrotus purpuratus)		
8.13	Wastewater Testing According to Dinnel and Stober	(1987)	using Red Sea Urchin
	(Strongylocentrotus franciscanus)		
8.14	Wastewater Testing According to Dinnel and Stober (Dendraster excentricus)	(1987)	using Sand Dollar
	(Dendraster excentricus)		
8.15	Wastewater Testing According to procedure E 724-8	9 (ASTM.	1989) using Pacific Oyster
	(Crassostrea gigas)		
8.16	Wastewater Testing According to procedure E 724-8	39 (ASTM.	1989) using California Bay Mussel
		-	
8.17	Uasteuater Testing According to Standard Methods	CADHA 1	080) using an alga
0.17	Wastewater Testing According to Standard Methods (Skeletonema costatum)	(AFIA, 1	707) using an argu
8.18	Wastewater Testing According to EPA/600/4-90/027		
0.10	Wastewater reseming According to EPA/000/4-70/02/	using ri	estinater and or rial the organismo
9	Physical Properties Testing of Hazardous Waste	(07-15-9	$\overline{\mathbf{p}}$
9.1	Ignitability by flashpoint determination (Title 2	פים ככ	66261.21)
	Corrosivity - pH determination (Title 22, CCR, 60	(241 22)	002011217
9.2	Corrosivity - Corrosivity towards steel (Title 22	201.22)	(2/4 22)
9.3	Corrosivity - Corrosivity towards steel (little 22	2, CCR, 6	0201.22)
9.4	Reactivity (Title 22, CCR, 66261.23)	• • • • • • • • • • • • • • • • • • • •	
10	Inorganic Chemistry and Toxic Chemical Elements	of Haza	rdous Waste
	A-A-1	40 -	ashal A
10.1	Antimony	10.7	Cobalt
	7040(N		7200()
	7041(06-06-86) Y		7201()
10.2	Arsenic	10.8	Copper
	7060(06-06-86) Y		7210()
	7061(N		7211()
10.7		10.0	
10.3	Barium 7080() N	10.9	Lead 7420(06-06-86)
	700U() N		7420(06-06-86)
	7081(N		
10.4	Beryllium	10.10	Mercury
	7090(N		7470()
	7091(N		7471(11-30-93)
10.5	Cadmium	10 11	Mol vhdenim
10.5	7130(N	10.11	7480()
	7131(N		7481()
10.6	Chromium, total	10.12	Nickel
	7190(N		7520()
	7903/		

CERTIFICATE NUMBER: 1573 EXPIRATION DATE: 7/31/97

10.13	Selenium		
	7740(06-06-86) Y	10.19	Cyanide
	7741(N		9010(06-06-86) Y
10.14	Silver	10.20	Fluoride
	7760(06-06-86) Y		300.0(11-30-93) ····································
	7761(N		340.1() N
10.15	Thallium		340.2(06-06-86) · · · · · · Y
	7840(N		340.3(N
	7841(06-06-86) Y	10.21	Sulfide
10.16	Vanadium		9030(06-06-86)Y
	7910(N	10.22	Total Organic Lead
	7911(N		(N
10.17	Zinc	10.23	EPA Method 6010(06-06-86) Y
	7950(N	10.24	EPA Method 6020(11-30-93) Y
	7951(N		
10.18	Chromium (VI)		
	7195(N		
	7196(06-06-86)Y		
	7197(N		
	7198(N		
	7 7 7		
11	Extraction Tests of Hazardous Waste (06-06-86	6)	
• • •	EXTINCTION TESTS OF HUZZA GOOD WASTE TOO OF OR	27	
11.1	California Waste Extraction Test (WET) (Title	22 CCB 66	261 100 Appendix II) Y
11.2	Extraction Procedure Toxicity	22, 000, 00	Y
	Toxicity Characteristic Leaching Procedure (TO	CLDS ALL CLA	, , , , , , , , , , , , , , , , , , ,
11.3	Toxicity Characteristic Leaching Procedure (TO	CLP) ALL CLE	vice Coly N
11.4	Toxicity Characteristic Leaching Procedure (10	CLP) Inorgan	nics only
11.5	Toxicity Characteristic Leaching Procedure (TO	CLP) EXTRACT	ables unity
11.6	Toxicity Characteristic Leaching Procedure (TO	CLP) Volatil	es uniy
			, , , , , , , , , , , , , , , , , , , ,
12	Organic Chemistry of Hazardous Waste (measur	ement by GC	/MS_combination)
	EPA Method 8240(02-05-87)		v
12.1	EPA Method 8240(02-05-87) EPA Method 8250()		······································
12.2	EPA Method 8250()		
12.3	EPA method 8270(02-05-87)		Υ
12.4	EPA Method 8280()		N
12.5	EPA Method 8290()		
12.6	EPA Method 8260(11-30-93)		ү
13	Organic Chemistry of Hazardous Waste (exclud	ing measure	ments by GC/MS combination)
13.1	EPA Method 8010(01-24-96) Y	13.13	EPA Method 8310(05-27-92) Y
13.2	EPA Method 8015(01-09-90) Y	13.14	
13.3	EPA Method 8020(06-06-86) Y	13.15	Total Petroleum Hydrocarbons
13.4	EPA Method 8030() N		(LUFT Manual) (06-06-86) Y
13.5	EPA Method 8040(N	13.16	EPA Method 8011(11-08-87) Y
13.6	EPA Method 8060(08-31-95) Y	13.17	' EPA Method 8021(-1-24-96) Y
13.7	EPA Method 8080(06-06-86) Y	13.18	B EPA Method 8070(N
13.8	EPA Method 8090() N	13.19	P EPA Method 8110() N
13.9	EPA Method 8100(N	13.20	EPA Method 8141(11-30-93) Y
13.10		13.21	
13.11			
	PA Method 8150(06-06-86) Y		
14	Bulk Asbestos Analysis ()		
14.1	1% or Greater Asbestos Concentrations (Title	22, CCR, 66	261.24(a)(2)(A))N
		,	
15	Substances Regulated Under the California Sa	afe Drinkin	g Water and Toxic Enforcement Act
	(Proposition 65) and Not Included in Other		
16	Wastewater Inorganic Chemistry, Nutrients a	nd Demand (07-15-91)
16.1	Acidity Y	16.1	3 Cyanide amenable to Chlorination Y
16.2		16.1	
16.3	_ '	16.1	
16.4		16.1	· · · · · · · · · · · · · · · · · · ·
16.5			
16.6		16.1	
		16.1	
16.7		16.1	
16.8		16.2	
16.9		16.2	urganic Carbon
16.1		16.2	2 Oxygen, Dissolved
16.1			
16.1	2 Cyanide Y		

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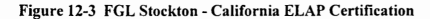
16.23 16.24 16.25 16.26	pH Y Phenols Y Phosphate, ortho Y Phosphorus, total Y	16.39 16.40 16.41 16.42	Surfactants (MBAS) Y Tannin and Lignin Y Turbidity Y Iron (Colorimetric Only) N
16.27	Potassium Y	16.43	Manganese (Colorimetric Only) N
16.28	Residue, Total Y	16.44	Total Recoverable
16.29	Residue, Filterable (TDS) Y		Petroleum Hydrocarbons Y
16.30	Residue, Nonfilterable (TSS) Y	16.45	Total Organic Halides Y
16.31	Residue, Settleable (SS) Y		
16.32	Residue, Volatile Y		
16.33	Silica Y Sodium Y		
16.34 16.35	Specific Conductance Y		
16.36	Sulfate Y		
16.37	Sulfide (includes total & soluble) - Y		
16.38	SulfiteY		
17	Toxic Chemical Elements in Wastewater (07		
17.1	Aluminum Y	17.18	Nicket Y Osmium N
17.2	Antimony ····· Y	17.19	Osmium N Palladium N
17.3	Arsenic Y Barium Y	17.20	Platinum N
17.4	Beryllium Y	17.21	Rhodium N
17.5 17.6	Cadmium Y	17.22 17.23	Ruthenium N
17.7	Chromium (VI)Y	17.24	Selenium Y
17.8	Chromium, total Y	17.25	Silver Y
17.9	CobaltY	17.26	Strontium Y
17.10	Copper Y	17.27	Thattium Y
17.11	Gold Y	17.28	Tin Y
17.12	Iridium N	17.29	Titanium Y
17.13	Iron Y	17.30	Vanadium Y
17.14	Lead Y	17.31	Zinc Y
17.15	Manganese ····· Y	17.32	EPA Method 200.7 Y
17.16	Mercury Y Molybdenum Y	17.33	EPA Method 200.8 Y
17.17	Molybdenum	17.34 17.35	Asbestos · N
		17.33	ASDESTOS TOTAL
18	Organic Chemistry of Wastewater (measure	ments by GC/MS	
18 18.1	EDA Mathod 674	nents by GC/MS	
	EPA Method 624	ments by GC/MS	
18.1 18.2 18.3	EPA Method 624 EPA Method 625 EPA Method 1613	ments by GC/MS	· \
18.1 18.2 18.3 18.4	EPA Method 624 EPA Method 625 EPA Method 1613	nents by GC/MS	·
18.1 18.2 18.3	EPA Method 624 EPA Method 625 EPA Method 1613	nents by GC/MS	· \
18.1 18.2 18.3 18.4	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625 EPA Method 613 Organic Chemistry of Wastewater (excludit	nents by GC/MS	by GC/MS combination) (07-15-91)
18.1 18.2 18.3 18.4 18.5	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 613 Organic Chemistry of Wastewater (excluding parts)	ments by GC/MS	by GC/MS combination) (07-15-91)
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18.1 18.2 18.3 18.4 18.5 19 19.1 19.2 19.3 19.4 19.5 19.7 20 20.1	EPA Method 625 EPA Method 1613 EPA Method 1625 EPA Method 613 Organic Chemistry of Wastewater (excluding Method 602 EPA Method 603 EPA Method 604 EPA Method 605 EPA Method 605 EPA Method 605 EPA Method 606 EPA Method 607 Inorganic Chemistry and Toxic Chemical E Processed Foods by One of the Following Method 606 Atomic Absorption Spectrophotometry Inductively Coupled Plasma Atomic Eminductively Coupled Plasma	ng measurements 19.8 19.9 19.10 19.11 19.12 19.13 Lements of Pest ethods ission Spectrop trometry ethods	by GC/MS combination) (07-15-91) EPA Method 608
18.1 18.2 18.3 18.4 18.5 19 19.1 19.2 19.3 19.4 19.5 19.7 20 20.1	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625 EPA Method 613 Organic Chemistry of Wastewater (excluding the second	ng measurements 19.8 19.9 19.10 19.11 19.12 19.13 Lements of Pest ethods ission Spectrop trometry ethods ission Spectrop trometry thods	by GC/MS_combination) (07-15-91) EPA Method 608

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20.4	Feed Products by One of the Following Methods Atomic Absorption Spectrophotometry Inductively Coupled Plasma Atomic Emission Spectrophotometry Inductively Coupled Plasma/Mass Spectrometry Colorimetry	N
21	Organic Chemistry of Pesticide Residues in Food (measurements by GC/MS) ()	
21.1 21.2 21.3 21.4	Gas Chromatographic/Mass Spectrometric Methods in Processed Foods	N
22	Organic Chemistry of Pesticide Residues in Food (Excluding Measurement by GC/MS Combination) ()	
22.1	Halogenated Compounds in Processed Foods by One of the Following Methods Gas Chromatography	N
22.2	Organophosphorous Compounds in Processed Foods by One of the Following Methods Gas Chromatography	N N
22.3	Carbamates in Processed Foods by One of the Following Methods Gas Chromatography	N
22.4	Halogenated Compounds in Raw Commodities by One of the Following Methods Gas Chromatography	N
22.5	Organophosphorous Compounds in Raw Commodities by One of the Following Methods Gas Chromatography	N
22.6	Carbamates in Raw Commodities by One of the Following Methods Gas Chromatography	N
22.7	Halogenated Compounds in Dairy Products by One of the Following Methods Gas Chromatography	N
22.8	Organophosphorous Compounds in Dairy Products by One of the Following Methods Gas Chromatography	N
22.9	Carbamates in Dairy Products by One of the Following Methods Gas Chromatography	
22.10	Gas Chromatography	. ,
22.11		-
22.12	Carbamates in Feed Products by One of the Following Methods Gas Chromatography	- 1 - 1

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Performance and System Audits



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ENVIRONMENTAL LABORATORY ACCREDITATION/REGISTRATION List of Approved Fields of Testing and Analytes

FGL Environmental 2500 Stagecoach Road Stockton, CA TELEPHONE No: (209) 942-0181
CALIFORNIA COUNTY: San Joaquin

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Microbiology of Drinking Water and Wastewater (07-03-91) Total Coliforms in Drinking Water by Multiple Tube Fermentation ------ Y 1.1 Fecal Coliforms/E. Coli in Drinking Water by MTF 1.2 Total Coliforms in Drinking Water by Membrane Filter Technics ------ N 1.3 Fecal Coliforms/E. Coli in Drinking Water by Membrane Filter Technics 1.4 Total Coliforms and E. Coli in Drinking Water by MMO-MUG -----Y 1.5 Total Coliforms in Drinking Water by Clark's Presence/Absence ------Y 1.6 Fecal Coliforms/E. Coli in Drinking Water by Clark's Presence/Absence ------Y 1.7 Heterotrophic Plate Count 1.8 Total Coliforms in Wastewater by Multiple Tube Fermentation 1.9 Fecal Coliforms in Wastewater by MTF 1.10 Total Coliforms in Wastewater by Membrane Filter Technics ----- N 1.11 Fecal Coliforms in Wastewater by Memorane Filter Technics ----- N 1.12 Fecal Streptococci or Enterococci by Multiple Tube Technics 1.13 Fecal Streptococci or Enterococci by Membrane Filter Technics ------ N 1.14 2 Inorganic Chemistry and Physical Properties of Drinking Water excluding Toxic Chemical Elements (07-03-91)Alkalinity ----- Y 2.12 Sulfate ----- Y 2.1 Calcium -----Y 2.13 Total Filterable Residue 2.2 Chloride -----Y and Conductivity ----- Y 2.3 Corrosivity ------ Y 2.14 Iron (Colorimetric Methods Only) ----- N 2.4 Fluoride -----Y 2.5 2.15 Manganese (Colorimetric Methods Only) - N Hardness ----- Y 2.16 Phosphate, ortho ----- Y 2.6 Magnesium ····· Y Silica (Colorimetric Methods Only) ---- Y 2.7 2.17 MBAS ----- Y 2.18 Cyanide ----- Y 2.8 Nitrate ----- Y 2.9 2.10 Sodium ----- Y 2.11 Analysis of Toxic Chemical Elements in Drinking Water (07-03-91) Arsenic ----- Y Silver ----- Y 3.1 3.11 Barium ----- Y Zinc ----- Y 3.2 3.12 Cadmium ----- Y Aluminum ------Y 3.3 3.13 Chromium, total Asbestos ----- N 3.4 3.14 Copper -----Y EPA Method 200.7 ----- Y 3.15 3.5 3.6 EPA Method 200.8 (Unregulated Elements 3.16 Lead ----- Y and Lead Only) ----- N 3.7 Manganese ------Y Antimony -----Y 3.8 3.17 Mercury ----- Y 3.9 3.18 Selenium ------Y Nickel -----Y 3.10 3.19 Thallium ----- Y 3.20 4 Organic Chemistry of Drinking Water (measurement by GC/MS combination) (07-03-91) EPA Method 501.3 ------Y 4.1 EPA Method 524.2 -----Y 4.2 EPA Method 525 ------4.3 EPA Method 513 4.4 5 Organic Chemistry of Drinking Water (excluding measurements by GC/MS combination) (07-03-91) EPA Method 501.1 ----- N EPA Method 547 ----- N 5.15 5.16 EPA Method 548 ----- N EPA Method 501.2 ----- Y EPA Method 549 ----- N EPA Method 502.1 ----- N 5.3 5.17 EPA Method 502.2 ----- Y EPA Method 550 ----- N 5.18 5.4 EPA Method 503.1 ----- Y
EPA Method 504 ----- Y EPA Method 550.1 ----- N 5.5 5.19 EPA Method 551 ----- N 5.6 5.20 EPA Method 505 ------ Y EPA Method 552 -----Y 5.7 5.21 EPA Method 506 5.8 EPA Method 508 ----- N 5.9 5.10 EPA Method 508A ----- N 5.11 EPA Method 510.1 ----- Y 5.12 EPA Method 515.1 ----- Y 5**.13** EPA Method 531.1 ------ N 5.14

Radiochemistry (-----) Gross Alpha and Beta Radiation ----- N 6.11 6.1 Gross Alpha by Co-precipitation ----- N 6.12 Total Radium ----- N 6.2 Radium 228 ----- N Radium 226 ----- N Radioactive Iodine ----- N 6.3 6.13 Uranium ------6.4 6.14 Gross Alpha & Beta in Hazardous Wastes -- N Radon 222 ----- N 6.5 Alpha Emitting Radium Isotopes 6.15 Radioactive Cesium -----in Haz. Wastes ----- N 6.6 Iodine 131 -----6.7 6.16 Radium 228 in Hazardous Wastes ----- N Radioactive Strontium ----- N 6.8 Tritium ----- N 6.9 Gamma and Photon Emitters ----- N 6.10 7 Shellfish Sanitation (-----) 7.1 Paralytic Shellfish Poison ------7.2 Domoic Acid ------N 7.3 8 Aquatic Toxicity Bioassays (-----) Hazardous Waste Aquatic Toxicity Bioassay (Title 22, CCR, 66261.24(a)(6)) ------ N 8.1 Wastewater Testing According to Kopperdahl (1976) using Freshwater Fish. ----- N 8.2 Wastewater Testing According to EPA/600/4-85/013 using Freshwater and/or Marine Organisms ------ N 8.3 8.4 8.5 Wastewater Testing by EPA Method 1003.0 8.6 Wastewater Testing by EPA Method 1006 N 8.7 Wastewater Testing by EPA Method 1007 -----N 8.8 Wastewater Testing by EPA Method 1009 8.9 8.10 Wastewater Testing According to Anderson, et. al. (1990) using Giant Kelp (Macrocystis pyrifera) -- N Wastewater Testing According to Anderson, et. al. (1990) using Red Abalone (Haliotus rufescens) --- N 8_11 Wastewater Testing According to Dinnel and Stober (1987) using Purple Sea Urchin 8.12 (Strongylocentrotus purpuratus) ------ N Wastewater Testing According to Dinnet and Stober (1987) using Red Sea Urchin 8.13 (Strongylocentrotus franciscanus) ------ N Wastewater Testing According to Dinnel and Stober (1987) using Sand Dollar 8.14 (Dendraster excentricus) ------ N Wastewater Testing According to procedure E 724-89 (ASTM, 1989) using Pacific Dyster 8.15 (Crassostrea gigas) ------ N Wastewater Testing According to procedure E 724-89 (ASTM, 1989) using California Bay Mussel 8.16 (Mytilus edulis) Wastewater Testing According to Standard Methods (APHA, 1989) using an alga-(Skeletonema costatum) ------ N Wastewater Testing According to EPA/600/4-90/027 using Freshwater and/or Marine Organisms ------ N 8.18 9 Physical Properties Testing of Hazardous Waste (11-09-93) Ignitability by Flashpoint determination (Title 22, CCR, 66261.21) ------ N 9.1 Corrosivity - pH determination (Title 22, CCR, 66261.22) ------Y 9.2 Corrosivity - Corrosivity towards steel (Title 22, CCR, 66261.22) ------ N 9.3 Reactivity (Title 22, CCR, 66261.23) ------N 9.4 Inorganic Chemistry and Toxic Chemical Elements of Hazardous Waste 10 Antimony 7200(----- N 7040(----- N 7201(----- N 7041(07-03-91) ----- Y 10.8 10.2 Arsenic Copper 7210(----- N 7060(07-03-91) ------ Y 7211(----- N 7061(----- N Lead 10.3 10.9 7080(----- N 7420(07-03-91) ----- Y 7421(----- N 7081(-----) ----- N 10.4 Beryllium 10.10 Mercury 7470(07-03-91) ----- Y 7090(----- N 7471(07-03-91) ----- Y 7091(----- N 10.11 Molybdenum Cadmium 10.5 7480(----- N 7130(----- N 7481(-----) ------ N 7131(----- N Nickel 10.6 Chromium, total 10.12 7520(-----) ----- N 7190(----- N 7191(----- N

10.13 Se	elenium		
7	7740(07-03-91) Y	10.19	Cyanide
7	7741(N		9010(N
10.14 Si	ilver	10.20	Fluoride
7	7760(07-03-91) Y		300.0(11-09-93) Y
7	7761(N		340.1(N
10.15	Thallium		340.2(07-03-91) Y
7	7840(N		340.3(N
	7841(07-03-91) Y	10.21	Sulfide
10.16	Vanadium		9030(N
	7910(N	10.22	Total Organic Lead
	7911(N	•	(07-03-91) Y
10.17	Zinc	10.23	EPA Method 6010(11-09-93) Y
	7950(N	10.24	EPA Method 6020(N
	7951(N		
	Chromium (VI)		
	7195(N		
	7196(07-03-91) Y		
	7197(N		
	7198(N		
	,		
11	Extraction Tests of Hazardous Waste (07-03-91)		
• •	EXTINCTION TESTS OF INDENTIONS MUSIC TOP US NIT		
11.1	California Waste Extraction Test (WET) (Title 22,	CCB 66	5261.100. Appendix II) Y
11.2	Extraction Procedure Toxicity		
11.3	Toxicity Characteristic Leaching Procedure (TCLP)	ALL CL	10000 U
11.4	Toxicity Characteristic Leaching Procedure (TCLP)	I Tooman	nice Only
	Toxicity Characteristic Leaching Procedure (TCLP)	Evenne	tables Only
11.5	Toxicity Characteristic Leaching Procedure (TCLP)) Extract	las Only
11.6	Toxicity Characteristic Leaching Procedure (TCLP)	volati	les unity
43	Onne in Observation of Management Marks (management	00	(MO
12	Organic Chemistry of Hazardous Waste (measureme	nt by Gu	/MS compination)
	EPA Method 8240(11-09-91)		- V
12.1	EPA Method 8250()	•••••	
12.2	EPA method 8270()	• • • • • • • • • • • • • • • • • • • •	
12.3	EPA method 82/0()		N
12.4	EPA Method 8280()		N
12.5	EPA Method 8290()		N
	EPA Method 8280()		N
12.5 12.6	EPA Method 8290()		N
12.5	EPA Method 8290()		N
12.5 12.6 13	EPA Method 8290()	measure	ements by GC/MS combination)
12.5 12.6 13	EPA Method 8290() EPA Method 8260(11-09-93) Organic Chemistry of Hazardous Waste (excluding EPA Method 8010(07-03-91)	measure	ements by GC/MS combination) S EPA Method 8310()
12.5 12.6 13 13.1 13.2	EPA Method 8290()	13.13 13.14	ments by GC/MS combination) S EPA Method 8310(No. EPA Method 632 (
12.5 12.6 13 13.1 13.2 13.3	EPA Method 8290() EPA Method 8260(11-09-93) Organic Chemistry of Hazardous Waste (excluding EPA Method 8010(07-03-91) EPA Method 8015(07-03-91) FPA Method 8020(07-03-91) FPA Method 8020(07-03-91)	measure	ements by GC/MS combination) S EPA Method 8310() N EPA Method 632 () N Total Petroleum Hydrocarbons
12.5 12.6 13 13.1 13.2 13.3 13.4	EPA Method 8290() EPA Method 8260(11-09-93) Organic Chemistry of Hazardous Waste (excluding EPA Method 8010(07-03-91) PPA Method 8015(07-03-91) EPA Method 8020(07-03-91) Y EPA Method 8030()	13.13 13.14 13.15	ments by GC/MS combination) EPA Method 8310() N EPA Method 632 () N Total Petroleum Hydrocarbons (LUFT Manual) (07-31-93)
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5	EPA Method 8290() EPA Method 8260(11-09-93) Organic Chemistry of Hazardous Waste (excluding EPA Method 8010(07-03-91)	13.13 13.14 13.15	### N Proceedings
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12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7	EPA Method 8290() EPA Method 8260(11-09-93) Organic Chemistry of Hazardous Waste (excluding EPA Method 8010(07-03-91)	13.13 13.14 13.15 13.16 13.17	N Y N Y N Y N Y N Y N Y N Y N N
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12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9	EPA Method 8290() EPA Method 8260(11-09-93) Organic Chemistry of Hazardous Waste (excluding EPA Method 8010(07-03-91) EPA Method 8015(07-03-91) EPA Method 8020(07-03-91) EPA Method 8030(13.13 13.14 13.15 13.16 13.17 13.18	N Y N Y N Y N Y N Y N Y N Y N N
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10	EPA Method 8290() EPA Method 8260(11-09-93) Organic Chemistry of Hazardous Waste (excluding EPA Method 8010(07-03-91) EPA Method 8015(07-03-91) EPA Method 8030() EPA Method 8040(07-03-91) EPA Method 8040(07-03-91) EPA Method 8060(13.13 13.14 13.15 13.16 13.17 13.18	ments by GC/MS combination) EPA Method 8310() Notal Petroleum Hydrocarbons (LUFT Manual) (07-31-93) Your York PA Method 8011(11-09-93) York PA Method 8021(01-25-96) York PA Method 8070() Notal Pethod 8070(
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11	EPA Method 8290() EPA Method 8260(11-09-93) Organic Chemistry of Hazardous Waste (excluding EPA Method 8010(07-03-91) EPA Method 8015(07-03-91) EPA Method 8030(07-03-91)	13.13 13.14 13.15 13.16 13.17 13.18	N Y N Y N Y N Y N Y N Y N Y N N
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11	EPA Method 8290() EPA Method 8260(11-09-93) Organic Chemistry of Hazardous Waste (excluding EPA Method 8010(07-03-91) EPA Method 8015(07-03-91) EPA Method 8030() EPA Method 8040(07-03-91) EPA Method 8040(07-03-91) EPA Method 8060(13.13 13.14 13.15 13.16 13.17 13.18	Nements by GC/MS combination Nements by GC/MS combination
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11	EPA Method 8290() EPA Method 8260(11-09-93) Organic Chemistry of Hazardous Waste (excluding) EPA Method 8010(07-03-91) Y EPA Method 8020(07-03-91) Y EPA Method 8030() N EPA Method 8040(07-03-91) Y EPA Method 8060() N EPA Method 8090() N EPA Method 8100() N EPA Method 8120() N EPA Method 8140() N EPA Method 8150(07-03-91) Y	13.13 13.14 13.15 13.16 13.17 13.18	Nements by GC/MS combination Nements by GC/MS combination
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11	EPA Method 8290() EPA Method 8260(11-09-93) Organic Chemistry of Hazardous Waste (excluding EPA Method 8010(07-03-91) EPA Method 8015(07-03-91) EPA Method 8030(07-03-91)	13.13 13.14 13.15 13.16 13.17 13.18	Nements by GC/MS combination Nements by GC/MS combination
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11	EPA Method 8290() EPA Method 8260(11-09-93) Organic Chemistry of Hazardous Waste (excluding EPA Method 8010(07-03-91) Y EPA Method 8020(07-03-91) Y EPA Method 8030() N EPA Method 8040(07-03-91) Y EPA Method 8060() N EPA Method 8090(07-03-91) Y EPA Method 8100() N EPA Method 8120() N EPA Method 8140() N EPA Method 8150(07-03-91) Y Bulk Asbestos Analysis (13.13 13.14 13.15 13.16 13.17 13.18 13.19 13.20	N Y N Y N Y N Y N Y N Y N Y N N
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11	EPA Method 8290() EPA Method 8260(11-09-93) Organic Chemistry of Hazardous Waste (excluding EPA Method 8010(07-03-91) Y EPA Method 8020(07-03-91) Y EPA Method 8030() N EPA Method 8040(07-03-91) Y EPA Method 8060() N EPA Method 8090(07-03-91) Y EPA Method 8100() N EPA Method 8120() N EPA Method 8140() N EPA Method 8150(07-03-91) Y Bulk Asbestos Analysis (13.13 13.14 13.15 13.16 13.17 13.18 13.19 13.20	N Y N Y N Y N Y N Y N Y N Y N N
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11 13.12	EPA Method 8290() EPA Method 8260(11-09-93) Organic Chemistry of Hazardous Waste (excluding) EPA Method 8010(07-03-91) Y EPA Method 8020(07-03-91) Y EPA Method 8030() N EPA Method 8040(07-03-91) Y EPA Method 8060() N EPA Method 8090(07-03-91) Y EPA Method 8100() N EPA Method 8120() N EPA Method 8140() N EPA Method 8150(07-03-91) Y Bulk Asbestos Analysis () Y 1% or Greater Asbestos Concentrations (Title 22,	13.13 13.14 13.15 13.16 13.17 13.18 13.19 13.20	N Y N Y N Y N Y N Y N N
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11 13.12	EPA Method 8290() EPA Method 8260(11-09-93) Organic Chemistry of Hazardous Waste (excluding EPA Method 8010(07-03-91) Y EPA Method 8020(07-03-91) Y EPA Method 8030() N EPA Method 8040(07-03-91) Y EPA Method 8060() N EPA Method 8090(07-03-91) Y EPA Method 8100() N EPA Method 8120() N EPA Method 8140() N EPA Method 8150(07-03-91) Y Bulk Asbestos Analysis (13.13 13.14 13.15 13.16 13.17 13.18 13.19 13.20	N Y N Y N Y N Y N Y N N
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11 13.12	EPA Method 8290() EPA Method 8260(11-09-93) Organic Chemistry of Hazardous Waste (excluding) EPA Method 8010(07-03-91) Y EPA Method 8020(07-03-91) Y EPA Method 8030() N EPA Method 8040(07-03-91) Y EPA Method 8060() N EPA Method 8090(07-03-91) Y EPA Method 8100() N EPA Method 8120() N EPA Method 8140() N EPA Method 8150(07-03-91) Y Bulk Asbestos Analysis () Y 1% or Greater Asbestos Concentrations (Title 22,	13.13 13.14 13.15 13.16 13.17 13.18 13.19 13.20 13.20	N Y N Y N Y N Y N Y N Y N Y N N
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11 13.12	EPA Method 8290() EPA Method 8260(11-09-93) EPA Method 8010(07-03-91)	13.13 13.14 13.15 13.16 13.17 13.18 13.19 13.20 13.20	N Y N Y N Y N Y N Y N Y N N
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11 13.12	EPA Method 8290() EPA Method 8260(11-09-93) Organic Chemistry of Hazardous Waste (excluding) EPA Method 8010(07-03-91) Y EPA Method 8020(07-03-91) Y EPA Method 8030() Y EPA Method 8040(07-03-91) Y EPA Method 8040(07-03-91) Y EPA Method 8080(07-03-91) Y EPA Method 8090() N EPA Method 8100() N EPA Method 8120() N EPA Method 8150(07-03-91) Y BA Method 8150(07-03-91) Y BULK Asbestos Analysis () Y Substances Regulated Under the California Safe	13.13 13.14 13.15 13.16 13.17 13.18 13.19 13.20 13.20	N Y N Y N Y N Y N Y N Y N N
12.5 12.6 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11 13.12	EPA Method 8290() EPA Method 8260(11-09-93) EPA Method 8010(07-03-91)	13.13 13.14 13.15 13.16 13.17 13.18 13.19 13.20 13.20	N Y
12.5 12.6 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11 13.12	EPA Method 8290() EPA Method 8260(11-09-93) EPA Method 8010(07-03-91)	13.13 13.14 13.15 13.16 13.17 13.18 13.19 13.20 13.20	N Y N Y N Y N Y N Y N Y N Y N N
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11 13.12	EPA Method 8290() EPA Method 8260(11-09-93) EPA Method 8010(07-03-91) EPA Method 8015(07-03-91) Y EPA Method 8020(07-03-91) Y EPA Method 8030() Y EPA Method 8040(07-03-91) Y EPA Method 8060() N EPA Method 8090() N EPA Method 8100() N EPA Method 8100() N EPA Method 8100() N EPA Method 8150(07-03-91) Y Bulk Asbestos Analysis () Y Bulk Asbestos Analysis (13.13 13.14 13.15 13.16 13.17 13.18 13.19 13.20 13.20	N Y N Y N Y N Y N Y N Y N Y N N
12.5 12.6 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11 13.12 14 14.1	EPA Method 8290() EPA Method 8260(11-09-93) EPA Method 8010(07-03-91)	13.13 13.14 13.15 13.16 13.17 13.18 13.19 13.20 13.20	N Y N Y N Y N Y N Y N Y N Y N N
12.5 12.6 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11 13.12 14 14.1	EPA Method 8290(13.13 13.14 13.15 13.16 13.17 13.18 13.19 13.20 13.20 13.20	N Y N Y N Y N Y N Y N Y N Y N N
12.5 12.6 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11 13.12 14 14.1	EPA Method 8290() EPA Method 8260(11-09-93) EPA Method 8010(07-03-91) Y EPA Method 8015(07-03-91) Y EPA Method 8020(07-03-91) Y EPA Method 8030() Y EPA Method 8040(07-03-91) Y EPA Method 8060() N EPA Method 8090() N EPA Method 8100() N EPA Method 8100() N EPA Method 8150(07-03-91) Y EPA Method 8150(07-03-91) Y BPA Method 8150(07-03-91) Y BULK Asbestos Analysis () Y Bulk Asbestos Analysis () Y Substances Regulated Under the California Safe (Proposition 65) and Not Included in Other Lis Wastewater Inorganic Chemistry, Nutrients and Included Chemistry Sutrients and Included Chemistry	13.13 13.14 13.15 13.16 13.17 13.18 13.19 13.20 13.20 13.20 13.20 13.20 13.20	N Y N Y N Y N Y N Y N Y N Y N N
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11 14.1 15	EPA Method 8290(13.13 13.14 13.15 13.16 13.17 13.18 13.19 13.20 13.20 13.20 13.20 13.20 13.20 13.20 13.20	N Y Y Y Y Y Y Y Y Y
12.5 12.6 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11 13.12 14 14.1 15	EPA Method 8290(13.13 13.14 13.15 13.16 13.17 13.18 13.20 13.20 13.20 13.20 13.20 13.20 13.20 13.20 13.20 13.20	N Y Y Y Y Y Y Y Y Y
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11 13.12 14 14.1 15	EPA Method 8290() EPA Method 8260(11-09-93) EPA Method 8010(07-03-91) Y EPA Method 8015(07-03-91) Y EPA Method 8020(07-03-91) Y EPA Method 8030() Y EPA Method 8040(07-03-91) Y EPA Method 8060() N EPA Method 8090() N EPA Method 8100() N EPA Method 8120(13.13 13.14 13.15 13.16 13.17 13.18 13.19 13.20 13.20 13.20 13.20 13.20 13.20 14.11 16.11 16.11 16.11 16.11	N Y N Y N Y N Y N Y N Y N Y N N
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11 13.12 14 14.1 15	EPA Method 8290() EPA Method 8260(11-09-93) EPA Method 8010(07-03-91) Y EPA Method 8015(07-03-91) Y EPA Method 8020(07-03-91) Y EPA Method 8030() N EPA Method 8040(07-03-91) Y EPA Method 8080(07-03-91) Y EPA Method 8090() N EPA Method 8100() N EPA Method 8120() N EPA Method 8150(07-03-91) Y Bulk Asbestos Analysis () Y Bulk Asbestos Analysis (13.13 13.14 13.15 13.16 13.17 13.18 13.19 13.20 13.20 13.20 13.20 13.20 13.20 13.20 13.20 13.10 16.1 16.1 16.1 16.1 16.1 16.1	N Y N Y N Y N Y N Y N Y N Y N N
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11 13.12 14 14.1 15	EPA Method 8290(13.13 13.14 13.15 13.16 13.17 13.18 13.19 13.20	N Y N Y N Y N Y N Y N Y N Y N N
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11 13.12 14 14.1 15	EPA Method 8290(13.13 13.14 13.15 13.16 13.17 13.18 13.19 13.20 13.20 13.20 13.20 13.20 13.20 13.20 13.20 13.10 16.1 16.1 16.1 16.1 16.1 16.1	N Y N Y N Y N Y N Y N Y N Y N N
12.5 12.6 13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 13.10 13.11 13.12 14 14.1 15	EPA Method 8290(13.13 13.14 13.15 13.16 13.17 13.18 13.19 13.20	N Y N Y N Y N Y N Y N Y N Y N N

16.23	рн у	16.39	Surfactants (MBAS)	,
16.24	Phenolsy		Tannin and Lignin	
16.25	Phosphate, ortho-	16.40	Tannin and Lightii	٧.
		16.41	Turbidity	
16.26	Phosphorus, total	16.42	Iron (Colorimetric Only)	١
16.27	Potassium Y	16.43	Manganese (Colorimetric Only)	١
16.28	Residue, Total Y	16.44	Total Recoverable .	
16,29	Residue, Filterable (TDS) Y		Petroleum Hydrocarbons	,
16.30	Residue, Nonfilterable (TSS) Y	16.45	Total Organic Halides	
16.31	Residue, Settleable (SS) Y	10.43	Total Organic natioes	•
16.32	Residue, Volatile Y			
16.33	Silica Y			
16.34	Sodium Y			
16.35	Specific Conductance Y			
16.36	Sulfate Y			
16.37				
	Sulfide (includes total & soluble) - Y			
16.38	Sulfite Y			
17	Toxic Chemical Elements in Wastewater (07-03-91)	Ĺ		
17.1	Aluminum ····· Y	17.18	Nickel N	
17.2	Antimony Y	17.19	Osmium Y	
17.3	Arsenic Y	17.20	Palladium N	
17.4	Barium N	17.21	Platinum N	
17.5	Beryllium N		Rhodium N	
	Cadmium Y	17.22	Knodium	
17.6	Cacini un	17.23	Ruthenium N	
17.7	Chromium (VI) Y	17.24	Selenium Y	
17.8	Chromium, total Y	17.25	Silver Y	
17.9	Cobalt N	17.26	Strontium N	
17.10	Copper N	17.27	Thallium Y	
17.11	Gold N	17.28	Tin Y	
	Iridium N		Titanium N	
17.12		17.29	i i tanium	
17.13	Iron N	17.30	Vanadium N	
17.14	Lead ····· Y	17.31	Zinc N	
17.15	Manganese N	17.32	EPA Method 200.7 Y	
17.16	Mercury Y	17.33	EPA Method 200.8 N	
17.17	Motybdenum N	17.34	DCP N	
		17 35	Achaetae N	
		17.35	Asbestos N	
18	Organic Chamistry of Usetquater (massuraments by			
18	Organic Chemistry of Wastewater (measurements b			
		y GC/MS	combination (07-03-91)	
18.1	EPA Method 624	y GC/MS	combination (07-03-91)	Y
18.1 18.2	EPA Method 624	y GC/MS	combination (07-03-91)	Y
18.1	EPA Method 624 EPA Method 625 EPA Method 1613	y GC/MS	combination (07-03-91)	YN
18.1 18.2	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625	y GC/MS	combination (07-03-91)	Y N N N N N
18.1 18.2 18.3	EPA Method 624 EPA Method 625 EPA Method 1613	y GC/MS	combination (07-03-91)	YN
18.1 18.2 18.3 18.4	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625	y GC/MS	combination (07-03-91)	Y N N N N N
18.1 18.2 18.3 18.4	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625	y GC/MS	combination (07-03-91)	Y N N N N N
18.1 18.2 18.3 18.4 18.5	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625 EPA Method 613 Organic Chemistry of Wastewater (excluding meas	y GC/MS	by GC/MS combination) (07-03-91)	Y N N N N N N N N
18.1 18.2 18.3 18.4 18.5	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601	y GC/MS	by GC/MS combination) (07-03-91)	Y N N N N N N N N N N N N N N N N N N N
18.1 18.2 18.3 18.4 18.5	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601 Y EPA Method 602	y GC/MS	by GC/MS combination) (07-03-91) EPA Method 608	Y N N N N
18.1 18.2 18.3 18.4 18.5 19	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601 Y EPA Method 602	urements 19.8 19.9	by GC/MS combination) (07-03-91) EPA Method 608	Y N N N N
18.1 18.2 18.3 18.4 18.5 19 19.1 19.2 19.3	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601 EPA Method 602 Y EPA Method 603	y GC/MS (by GC/MS combination) (07-03-91) EPA Method 608	Y N N N N
18.1 18.2 18.3 18.4 18.5 19 19.1 19.2 19.3 19.4	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601 EPA Method 602 EPA Method 603 EPA Method 603 EPA Method 604	19.8 19.9 19.10	by GC/MS combination) (07-03-91) EPA Method 608	Y N N N N
18.1 18.2 18.3 18.4 18.5 19 19.1 19.2 19.3 19.4 19.5	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601 EPA Method 602 EPA Method 603 EPA Method 604 EPA Method 604 EPA Method 605 N	19.8 19.9 19.10 19.11	by GC/MS combination) (07-03-91) EPA Method 608	Y N N N N
18.1 18.2 18.3 18.4 18.5 19 19.1 19.2 19.3 19.4 19.5 19.6	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601 EPA Method 602 EPA Method 603 EPA Method 604 EPA Method 604 EPA Method 605 EPA Method 605 EPA Method 606	19.8 19.9 19.10	by GC/MS combination) (07-03-91) EPA Method 608	Y N N N N
18.1 18.2 18.3 18.4 18.5 19 19.1 19.2 19.3 19.4 19.5	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601 EPA Method 602 EPA Method 603 EPA Method 604 EPA Method 604 EPA Method 605 N	19.8 19.9 19.10 19.11	by GC/MS combination) (07-03-91) EPA Method 608	Y N N N N
18.1 18.2 18.3 18.4 18.5 19 19.1 19.2 19.3 19.4 19.5 19.6	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601 EPA Method 602 EPA Method 603 EPA Method 604 EPA Method 604 EPA Method 605 EPA Method 605 EPA Method 606	19.8 19.9 19.10 19.11	by GC/MS combination) (07-03-91) EPA Method 608	Y N N N N
18.1 18.2 18.3 18.4 18.5 19 19.1 19.2 19.3 19.4 19.5 19.6 19.7	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601 EPA Method 602 EPA Method 603 EPA Method 603 EPA Method 604 EPA Method 605 EPA Method 605 EPA Method 605 EPA Method 606 EPA Method 606 N EPA Method 607 N EPA Method 607	19.8 19.9 19.10 19.11 19.12	by GC/MS combination) (07-03-91) EPA Method 608	Y N N N N
18.1 18.2 18.3 18.4 18.5 19 19.1 19.2 19.3 19.4 19.5 19.6	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601 EPA Method 602 EPA Method 603 EPA Method 604 EPA Method 604 EPA Method 605 EPA Method 605 EPA Method 606	19.8 19.9 19.10 19.11 19.12	by GC/MS combination) (07-03-91) EPA Method 608	Y N N N N
18.1 18.2 18.3 18.4 18.5 19 19.1 19.2 19.3 19.4 19.5 19.6 19.7	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601 EPA Method 602 EPA Method 603 EPA Method 603 EPA Method 604 EPA Method 605 EPA Method 605 EPA Method 605 EPA Method 606 EPA Method 606 N EPA Method 607 N EPA Method 607	19.8 19.9 19.10 19.11 19.12	by GC/MS combination) (07-03-91) EPA Method 608	Y N N N N
18.1 18.2 18.3 18.4 18.5 19 19.1 19.2 19.3 19.4 19.5 19.6 19.7	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601 EPA Method 602 EPA Method 603 EPA Method 604 EPA Method 604 EPA Method 605 EPA Method 606 EPA Method 607 Inorganic Chemistry and Toxic Chemical Elements Processed Foods by One of the Following Methods	19.8 19.9 19.10 19.11 19.12 19.13	by GC/MS combination) (07-03-91) EPA Method 608	Y
18.1 18.2 18.3 18.4 18.5 19 19.1 19.2 19.3 19.4 19.5 19.7	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601 EPA Method 602 EPA Method 603 EPA Method 604 EPA Method 604 EPA Method 605 EPA Method 606 EPA Method 607 Inorganic Chemistry and Toxic Chemical Elements Processed Foods by One of the Following Methods Atomic Absorption Spectrophotometry	19.8 19.9 19.10 19.11 19.12 19.13	by GC/MS combination) (07-03-91) EPA Method 608	Y N N N N N N N N N N N N N N N N N N N
18.1 18.2 18.3 18.4 18.5 19 19.1 19.2 19.3 19.4 19.5 19.7	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601 EPA Method 602 EPA Method 603 EPA Method 604 EPA Method 605 EPA Method 605 EPA Method 606 EPA Method 607 Inorganic Chemistry and Toxic Chemical Elements Processed Foods by One of the Following Methods Atomic Absorption Spectrophotometry Inductively Coupled Plasma Atomic Emission	19.8 19.9 19.10 19.11 19.13 of Pesti	by GC/MS combination) (07-03-91) EPA Method 608	Y N N N N
18.1 18.2 18.3 18.4 18.5 19 19.1 19.2 19.3 19.4 19.5 19.7	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601 EPA Method 602 EPA Method 603 EPA Method 604 EPA Method 605 EPA Method 605 EPA Method 606 EPA Method 607 Inorganic Chemistry and Toxic Chemical Elements Processed Foods by One of the Following Methods Atomic Absorption Spectrophotometry Inductively Coupled Plasma Atomic Emission	19.8 19.9 19.10 19.11 19.13 of Pesti	by GC/MS combination) (07-03-91) EPA Method 608	Y N N N N
18.1 18.2 18.3 18.4 18.5 19 19.1 19.2 19.3 19.4 19.5 19.7	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601 EPA Method 602 EPA Method 603 EPA Method 604 EPA Method 605 EPA Method 606 EPA Method 607 Inorganic Chemistry and Toxic Chemical Elements Processed Foods by One of the Following Methods Atomic Absorption Spectrophotometry Inductively Coupled Plasma Atomic Emission Inductively Coupled Plasma/Mass Spectrometry	19.8 19.9 19.10 19.11 19.13 of Pesti	by GC/MS combination) (07-03-91) EPA Method 608	Y N N N N N N N N N N N N N N N N N N N
18.1 18.2 18.3 18.4 18.5 19 19.1 19.2 19.3 19.4 19.5 19.6 19.7	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601 EPA Method 602 EPA Method 603 EPA Method 604 EPA Method 605 EPA Method 606 EPA Method 607 Inorganic Chemistry and Toxic Chemical Elements Processed Foods by One of the Following Methods Atomic Absorption Spectrophotometry Inductively Coupled Plasma Atomic Emission Inductively Coupled Plasma Atomic Emission Inductively Coupled Plasma/Mass Spectrometry Colorimetry	19.8 19.9 19.10 19.11 19.13 of Pesti	by GC/MS combination) (07-03-91) EPA Method 608	Y N N N N N N N N N N N N N N N N N N N
18.1 18.2 18.3 18.4 18.5 19 19.1 19.2 19.3 19.4 19.5 19.7	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601 EPA Method 602 EPA Method 603 EPA Method 604 EPA Method 605 EPA Method 606 EPA Method 606 EPA Method 607 Inorganic Chemistry and Toxic Chemical Elements Processed Foods by One of the Following Methods Atomic Absorption Spectrophotometry Inductively Coupled Plasma Atomic Emission Inductively Coupled Plasma Atomic Emission Inductively Coupled Plasma Atomic Emission Inductively Coupled Plasma/Mass Spectrometr Colorimetry Ray Commodities by One of the Following Methods	19.8 19.9 19.10 19.11 19.13 of Pesti	by GC/MS combination) (07-03-91) EPA Method 608	Y N N N N N N N N N N N N N N N N N N N
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18.1 18.2 18.3 18.4 18.5 19 19.1 19.2 19.3 19.4 19.5 19.6 19.7	EPA Method 624 EPA Method 625 EPA Method 1613 EPA Method 1625 EPA Method 613 Organic Chemistry of Wastewater (excluding meas) EPA Method 601 EPA Method 602 EPA Method 603 EPA Method 604 EPA Method 605 EPA Method 606 EPA Method 606 EPA Method 607 Inorganic Chemistry and Toxic Chemical Elements Processed Foods by One of the Following Methods Atomic Absorption Spectrophotometry Inductively Coupled Plasma/Mass Spectrometric Colorimetry Raw Commodities by One of the Following Methods Atomic Absorption Spectrophotometry Inductively Coupled Plasma/Mass Spectrometric Colorimetric Colorimetric Colorimetric Colorimetric Dairy Rooderts by One of the Following Methods	19.8 19.9 19.10 19.11 19.13 of Pesti	by GC/MS combination) (07-03-91) EPA Method 608	Y N N N N
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20.4	Feed Products by One of the Following Methods Atomic Absorption Spectrophotometry Inductively Coupled Plasma Atomic Emission Spectrophotometry Inductively Coupled Plasma/Mass Spectrometry Colorimetry	N
21	Organic Chemistry of Pesticide Residues in Food (measurements by GC/MS) ()	
21.1	Gas Chromatographic/Mass Spectrometric Methods in Processed Foods	N
21.2	Gas Chromatographic/Mass Spectrometric Methods in Ray Commodities	N
21.3	Gas Chromatographic/Mass Spectrometric Methods in Dairy Products	N
21.4	Gas Chromatographic/Mass Spectrometric Methods in Feed Products	١
22	Organic Chemistry of Pesticide Residues in Food (Excluding Measurement by GC/MS Combination)	
22.1	Halogenated Compounds in Processed Foods by One of the Following Methods	
	Gas Chromatography	N
	High Pressure Liquid Chromatography	N
	Liquid Chromatography/Mass Spectrometry	N
22.2	Organophosphorous Compounds in Processed Foods by One of the Following Methods	
	Gas Chromatography	N
	High Pressure Liquid Chromatography	N
	Liquid Chromatography/Mass Spectrometry	N
22.3	Carbamates in Processed Foods by One of the Following Methods	
	Gas Chromatography	N
	High Pressure Liquid Chromatography	N
	Liquid Chromatography/Mass Spectrometry	N
22.4	Halogenated Compounds in Raw Commodities by One of the Following Methods	
	Gas Chromatography	N
	High Pressure Liquid Chromatography	N
	Liquid Chromatography/Mass Spectrometry	N
22.5	Organophosphorous Compounds in Raw Commodities by One of the Following Methods Gas Chromatography	
	Gas Chromatography	N
	High Pressure Liquid Chromatography	N
77.	Liquid Chromatography/Mass Spectrometry	N
22.6	Carbamates in Raw Commodities by One of the Following Methods Gas Chromatography	
	High Pressure Liquid Chromatography	N
	Liquid Chromatography/Mass Spectrometry	N
22.7	Halogenated Compounds in Dairy Products by One of the Following Methods	r
22.1	Gas Chromatography	L
	High Pressure Liquid Chromatography	
	Liquid Chromatography/Mass Spectrometry	
22.8	Organophosphorous Compounds in Dairy Products by One of the Following Methods	
	Gas Chromatography	
	High Pressure Liquid Chromatography	
	Liquid Chromatography/Mass Spectrometry	,
22.9	Carbamates in Dairy Products by One of the following Methods	
	Gas Chromatography	•
	High Pressure Liquid Chromatography	ł
	Liquid Chromatography/Mass Spectrometry	į
22.10	Halogenated Compounds in Feed Products by One of the Following Methods	
	Gas Chromatography	ł
	High Pressure Liquid Chromatography	١
	Liquid Chromatography/Mass Spectrometry	1
22.11	Organophosphorous Compounds in Feed Products by One of the Following Methods	
	Gas Chromatography	-
	High Pressure Liquid Chromatography	
22 42	Liquid Chromatography/Mass Spectrometry	
22.12	Carbamates in Feed Products by One of the Following Methods Gas Chromatography	
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	High Pressure Liquid Chromatography	
	Liquid Chromatography/Mass Spectrometry	

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Performance and System Audits



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State of Utah

Michael O. Leavitt

Charles and

Rod L. Detit Faculty Opening

Charles D. Brokopp. Dr. P.H.

Bureau of Laboratory Improvement

- North Medical Drive 3 1. acc + 19. com 84110-1105 # iephone; 801 584-8469 .gesol 584-850

JAN 05 1996

DARRELL H NELSON FGL ENVIRONMENTAL PO BOX 272 SANTA PAULA CA 93061-0272

Certificate No.: E-278

Account No: 8056590910

On the basis of your most recent audit results, the laboratory listed is hereby certified for environmental monitoring under the Resource Conservation and Recovery Act and authorized to perform the following analytes, or groups of analytes by method:

METALS

ALUMINUM 6020

ANTIMONY 6020

ANTIMONY 7041

ARSENIC 6020

ARSENIC 7060

BARIUM 6010

BARIUM 6020

BERYLLIUM 6010

BERYLLIUM 6020

CADMIUM 6010

CADMIUM 6020

CHROMIUM 6010

CHROMIUM 6020

CHROMIUM, HEXAVALENT 7196

COBALT 6010

COBALT 6020

COPPER 6010 COPPER 6020

LEAD 6020

LEAD 7420

MANGANESE 6020

MERCURY 7470

MERCURY 7471

MOLYBDENUM 6010

NICKEL 6010

NICKEL 6020

SELENIUM 7741

SILVER 6010

SILVER 6020

THALLIUM 6020

THALLIUM 7841

VANADIUM 6010 ZINC 6010

ZINC 6020

MISCELLANEOUS

CYANIDE TOTAL/AMENABLE 9010

IGNITABILITY 1020

ORGANOCHL PEST 8080

ORGANOPHOSPHORUS PEST 8140

ORGANOPHOSPHORUS PEST 8141

PAH 8310

PAINT FILTER LIQUID TST 9095

PCB 8080

PH 9045

REACTIVITY SEC 2.1.3

REACTIVITY SEC 8.3

SEMIVOLATILES 8270

TCLP METAL 1311

TCLP VOLATILE 1311

TOTAL ORGANIC HALIDES 9020

VOLATILES 8240

VOLATILES 8260

RADIOLOGICS

GROSS ALPHA & BETA 9310

RADIUM 228 - 9320



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* of FallSY DEVI-

State of Utah

Michael O. Leavitt Covernor Rod L. Betit

Executive Director

Charles D. Brokopp, Dr. P.H.

Director

Bureau of Laboratory Improvement

46 North Medical Drive Sait Lake City, Utah 84113-1105 Telephone: 801 584-8469 Fax: 801 (584-8501)

JAN 05 1996

DARRELL H NELSON FGL ENVIRONMENTAL PO BOX 272 SANTA PAULA CA 93061-0272

Certificate No.: E-278

Account No:

8056590910

On the basis of your most recent audit results, the laboratory listed is hereby certified for environmental monitoring under the Safe Drinking Water Act and authorized to perform the following analytes, or groups of analytes by method:

RADIOLOGICS

CESIUM-134 GROSS ALPHA GROSS BETA RADIUM - 226

RADIUM-228 STRONTIUM - 89/90 TOTAL RADIUM TRITIUM URANIUM

This laboratory's certification is effective NOV 30 1995 through NOV 30 1997.

The analytes for which a laboratory is authorized to perform at any given time will be those indicated in the most recent certificate letter. Please review for completeness and accuracy. Any discrepancies must be documented and notice received by this Bureau within 15 days of receipt. Copies of this letter will be on file in (1) the Bureau of Laboratory Improvement, Division of Laboratory Services and (2) in the Department of Environmental Quality. The certification will be recalled in the event that your Laboratory's certification is revoked.

Respectfully,

Charles Brokopp, Dr. P.H.

Director

Richard Denton - DEQ (cc. electronic) Kenneth Bousfield - DEQ (cc. electronic) U.S. EPA Region VIII QAO

Management Services Coordinator (cc. electronic)

This laboratory's certification is effective JAN 23 1996 through NOV 30 1997.

The analytes for which a laboratory is authorized to perform at any given time will be those indicated in the most recent certificate letter. Please review for completeness and accuracy. Any discrepancies must be documented and notice received by this bureau within 15 days of receipt. Copies of this letter will be on file in (1) the Bureau of Laboratory Improvement, Division of Laboratory Services and (2) in the Department of Environmental Quality. The certification will be recalled in the event that your Laboratory's certification is revoked.

Respectfully,

Charles Brokopp, Dr. P.H.

2. Broken

Director

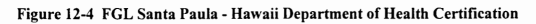
cc. Richard Denton - DEQ (cc. electronic)
 Dennis Downs - DEQ (cc. electronic)

U.S. EPA Region VIII QAO

Management Services Coordinator (cc. electronic)

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Performance and System Audits



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LAWRENCE MIKE

STATE OF HAWAII DEPARTMENT OF HEALTH

In reply, please refer to: File: SLD

2725 WAIMANO HOME ROAD PEARL CITY, HAWAII 96782-1496

It is recommended that data from the following laboratory be accepted for drinking water analyses for regulatory purposes for the contaminants listed until <u>February 28, 1997</u>.

PGL ENVIRONMENTAL

853 Corporation Street
Santa Paula, California 93061

Denis D. Barry, Marketing Director (805) 659-0910

INORGANICS	ORGANICS	VOLATILES
Antimony	Alachlor	THM
Arsenic	Atrazine	VOC
Barium	Chlordane	Vinyl Chloride
Beryllium	Endrin	•
Cadmium	Heptachlor	EDB
Chromium	Heptachlor Epoxide	DBCP
Copper	Hexachlorobenzene	
Lead	Hexachlorocyclopentadiene	
Hercury	Lindane	
Nickel	Methoxychlor	
Selenium	Simazine	OTHER
Thallium	Toxaphene	
	•	Residual Chlorine
Nitrate	λldicarb	TDS
Fluoride	Aldicarb Sulfone	Calcium
	Aldicarb Sulfoxide	рĦ
	Carbofuran	λlkalinity
ORGANICS	Oxamyl	Corrosivity
Endothall		Sodium
Glyphosate	2,4-D	Sulfate
Diquat	Silvex	
Bis-2-ethylhexyl Adipate	Pentachlorophenol	
Bis-2-ethylhexyl Phthalate	Dalapon	
Benzo(a)pyrene	Dinoseb	
	Picloram	
non describing house		

PCB as decachlorobiphenyl

RECOMMENDED:

Jodi Nakamura

Certification Officer

Date

2/13/96

Vernon K. Miyamoto, Ph.D.

APPROVED:

Date

Chief, State Laboratories Division

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Section No: 13 Page: 1 of 5 Revision No: 2.2 Date: May 9, 1995

Preventive Maintenance

13.1 Maintenance and Repair of Instruments

Routine maintenance of equipment is performed by the analyst when appropriate. The department supervisor must be notified immediately if any sign of serious malfunction occurs in any instrument so that he can decide if a qualified serviceman should be consulted. If warranted, instrument repair and calibration is performed by qualified service technicians (usually service representatives of the instrument manufacturer). A record containing the date, the analyst or service engineer, symptoms of the problem, description of the repair, and comments concerning the repair is also kept. Figures 13-1 and 13-2 give examples of the logbook cover and logbook record.

13.2 Refrigerator and Oven Temperature Monitoring

All refrigerators and ovens are monitored daily to confirm proper operation. Refrigerators and ovens which do not have an NIST certified thermometer are checked for accuracy using an NIST reference thermometer.

13.3 Balance Monitoring

All balances are monitored on a monthly basis to verify proper operation and accuracy. The weights used for verification are calibrated to class M2 tolerances. The weights are certified on an annual basis.

13.4 Good Laboratory Practices

Good laboratory practices are followed to prevent contamination of samples and standards. This includes the careful cleaning of glassware, and the use of disposable labware and containers when practical. Sample containers are monitored for contamination when received, according to lot number and proposed use.

The bacteriology water is monitored for suitability annually. Standard plate count, electrical conductivity and residual chlorine are checked monthly. Heavy metals (including lead, cadmium, chromium, copper, nickel and zinc) are checked annually.

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Figure 13-1 Repair/Maintenance Logbook Cover

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Analytical Chemists

Repair/Maintenance Log Book

Manuíacturer:
Model No
Serial No.
Instrument ID:

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Figure 13-1 Repair/Maintenance Logbook Record

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Activities Log		
Date:		
Analyst/Service Eng.:		
Symptoms:		
Repair/Maintenance:		
Comments:		

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Section No: 14 Page: 1 of 3 Revision No: 2.2 Date: May 9, 1995

Specific Routine Procedures used to assess Data Precision, Accuracy, and Completeness

Before analytical data can be used, it is necessary to determine the suitability of the data for a given purpose. The characteristics used to determine data suitability are precision, accuracy, and completeness. FGL Environmental deterimines these characteristics by using specific procedures, which are detailed in the following sections.

14.1 Precision

Precision is the measure of how closely replicate analyses agree. FGL Environmental uses Relative Percent Difference (RPD) to measure between duplicate analyses.

Precision is monitored for nearly all methods by RPD's plotted on control charts. The mean RPD +/- 2 standard deviations are the warning limits, and the mean RPD +/- 3 standard deviation are the control limits. To assess precision, FGL Environmental uses the following on a regular basis:

- (1) Duplicate samples
- (2) Duplicate Matrix Spikes
- (3) Control Charts

14.1.1 Precision Calculation

The RPD of duplicate samples is an absolute value from the following calculation:

(First Sample Value - Second Sample Value) X 100

(First Sample Value + Second Sample Value) / 2

14.2 Accuracy

Accuracy measures the deviation of the analytical value from the "true" or known value. The true value for field samples are never known, so accuracy measurements are made on the analysis of QC samples analyzed with field samples.

Accuracy is monitored for nearly all methods by percent recoveries plotted on control charts. The mean recovery +/- 2 standard deviation are the warning limits, and the mean recovery +/- 3 standard deviation are the control limits. To assess accuracy, FGL Environmental uses the following on a regular basis:

- (1) Initial Calibration Verification (ICV)
- (2) Continuing Calibration Verification (CCV)
- (3) Laboratory Control Samples (LCS)

FGL uses ICV and CCV to verify control for those analyses without significant sample preparation and a combination of ICV, CCV and LCS to verify control for those analyses which require significant sample preparation.

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Specific Routine Procedures used to assess Data Precision, Accuracy, and Completeness

14.2.1 Accuracy Calculations

Percent recoveries are calculated as follows (identical units would be used through each calculation):

ICV, CCV and LCS percent recoveries:

value found X 100

true value

14.3 Sample Matrix Effects

Sample matrix effect calculations measure the impact of the matrix on the analytical results. Matrix effects are control charted for the major matrix groups such as potable water, wastewater, solids, filters etc. However, FGL does not use the results for verifying analytical control.

Matrix spike and matrix spike duplicate results are monitored for nearly all methods and surrogate spike results are monitored for most organic methods by percent recoveries plotted on control charts. The mean recovery +/- 2 standard deviation are the warning limits, and the mean recovery +/- 3 standard deviation are the control limits.

14.3.1 Sample Matrix Effect Calculations

Percent recoveries are calculated as follows (identical units would be used through each calculation):

Spike percent recoveries:

(spiked sample result - sample result) X 100

spike amount added

14.4 Completeness

Completeness is defined by QAMS-005/80 as - a measure of the amount of valid data obtained from a measurement system compared to the amount that was expected to be obtained under correct normal conditions.

By this definition the influence of the laboratory on completeness involves three areas:

- 1) appropriate sample handling and storage,
- 2) conformance to holding time requirements and
- 3) data validity as measured by meeting acceptance criteria for the quality control parameters.

We do not track completeness as a measurable form at this point. We do strive to provide data packages that are 100% complete and give explanations when there are deficiencies.

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Specific Routine Procedures used to assess Data Precision, Accuracy, and Completeness

14.5 Control Charts

FGL is in the process of formalizing the use of quality assurance data. Many EPA and standard methods do not have formal quality assurance objectives. Also, EPA does not have criteria for dropping outliers from control charts. Below is an outline of the approach FGL is taking to provide for these deficiences.

14.5.1 Control Chart Limits

FGLwill be using several different control limits. The following is a list of limits with definitions:

- a) EPA limit limit specified in the method and required by EPA
- b) FGL control chart fixed limit used for CCV and LCS acceptance criteria. If an analysis or procedure does not have limits specified by the EPA the limits will be generated by control charts using a 99% confidence interval (3s). These limits will then be set and not allowed to be adjusted unless the control charts show a significant change in the method i.e. narrowing of 50%.
- c) FGL control chart variable limit used for BS, MS and RPD acceptance criteria. If an analysis or procedure does not have limits specified by the EPA the limits will be generated by control charts using a 99% confidence interval (3s) and perhaps a 95% confidence interval (2s), if a warning limit is deemed useful. These limits will then be allowed to fluctuate based on control chart findings.
- d) Global 3s control chart limits based on all points in the control chart less those points that are dropped based on criteria in section 2.
- e) Trend 3s control chart limits based on the most recent 30 points in the control chart less those points dropped based on criteria in section 2.

14.5.2 Criteria for Dropping Outliers

- a) RPD or % Rec. point drop low spike concentration the RPD or % rec. point may be dropped if the spike concentration is less than MAV times the sample concentration. i.e. if the sample concentration is 2.5 mg/l and the MAV is 15%, a spike of 0.374 mg/L or lower would be rejected as too low and a spike of 0.375 or greater would be maintained.
- b) RPD point drop low duplicate concentration the RPD point may be dropped if the sample concentration is lower than the DLR times the MAV. i.e. if the DLR is 0.2 mg/L and the MAV is 15%, an average sample concentration (for the duplicates) of 2.9 or lower would be rejected and an average sample concentration of 3.0 or above would be maintain.
- c) RPD or % Rec. outliers prior to setting the limits the points are reviewed. The point may be dropped if it fails the following test: If the control limits are shifted by 30% of the limit (i.e. 85 115% changing to 89.5 110.5%) or more by any single point then the point is considered an outlier and dropped.

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Date: October 3, 1994

Corrective Actions

Corrective actions are necessary when trends of more than one out-of- control situations occur. Corrective action reports are used to document the corrections made. Corrective actions are not normally used in isolated out-of-control situations that have routine explanations as the data in these situations must be resolved before continuing and reporting analyses.

15.1 Corrective Action Reports

Each work area has a corrective action report logbook. When corrective actions are necessary, a corrective action report form (figure 15-1) is filled out identifying the analyst, date, method, client and lab number (if applicable), QA batch number (if applicable) problems encountered, investigation and proposed corrective actions. After implementing the actions another entry is required to verify that the problem was solved. This process may need to be repeated in some situations. The reports are on record and will be included in a project data package if that is required by the project plan. An example of a corrective action report form is shown in figure 15-1.

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Figure 15-1 Corrective Action Report Form

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Corrective Actions

Corrective Action Report Form		
Method:		
Problem Assigned To:		
Problem encountered:		
		100 (1.00)
		·
Cause of the problem:		
A-10-1		
Corrective action:		
-		
Character of at		
Closure of Investigation:		
Performed by:		
Verified by:	Date:	

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Quality Assurance Reports to Management

In order to insure that the Quality Assurance program at the lab maintains a high profile, there are several mechanisms in place which insure the QA information is routinely conveyed to laboratory management. This includes a formal monthly QA inspection summary report, reports on internal and external PE samples and summary reports for external system audits.

16.1 Monthly QA Non-conformance Summary Reports

This report contains all the new non-conformance items, previously uncompleted non-conformance items and finally all non-conformance items completed. Dates are recorded when the non-conformance was found and when it was completed. This report is given to all supervisors and managers. With these steps an ongoing quantitative assessment of the analytical system is provided.

16.2 Annual QA Summary Reports

At the end of each year the QA director compiles all the quality assurance information for the previous year and provides a summary report to management. This report includes the number of non-conformances found throughout the year, trending information on the non-conformances, performance evaluation study summaries and evaluations of whether quality assurance inspection schedules were maintained.

16.3 Performance Evaluation Corrective Action Reports

Evaluations of any failures on external PE samples are outlined by department supervisors and prepared by the QA Director or Officer for certifying agencies. Copies are given to the department supervisors and Lab Director.

16.4 External System Audit Summary Reports

After debriefing by the auditors a summary report is prepared by the QA Director or Officer for the supervisors and Lab Director. Rather than waiting for an audit report, this initiates corrective actions for any non-conformance items promptly.

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Figure 16-1 FGL Monthly QA Non-Comformance Summary Report

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Quality Assurance Reports to Management

FGL Monthly QA Non-Comformance Summary Report

Current Non-conformance Items			
Non-Comformance Item	Person Resp.	Date Found	
Company I No.			
Corrected Non-comformance Items			
Non-Comformance Item	Person Resp.	Date Found	Date Comp.
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FGL Environmental Instrument and Equipment List

LAB	DIVISION	INSTRUMENT DESCRIPTION	SERIAL NUMBER	INST. TYPE	INST. ID	INV. CONTROL	REMARKS
SP	Organic	Hewlett Packard GC/MS 5890II/5972 with 7673A autosampler	3310A49023	GC/M\$	202	26-93-0046	
SP	Organic	Hewlett Packard GC/MS 589011/5972 with LSC2000/ALS2016 autosampler	3336A52456	GC/MS	204	26-94-0028	
SP	Organic	Hewlett Packard GC/MS 589011/5971 with 7673A autosampler	3203A41750	GC/MS	201	26-92-0036	
SP	Organic	Hewlett Packard GC/MS 589011/5971 with LSC2000/ALS2016 autosampler	2905A12089	GC/MS	205	26-95-0056-349	
SP	Organic	Hewlett Packard GC/MS 5890H/5970 with LSC2000/ALS2016 autosampler	3004A12738	GC/MS	206	26-95-0076-369	
SP	Organic	Varian 3400/Finnigan XL 50 with LSC2000/ALS2016/ALS2032 autosampler	IN001137	GC/MS	203	26-91-0025	
SP	Organic	Hewlett Packard GC 5890 with ECD + ECD detectors and 7673A autosampler	2750A15955	GC	202	26-95-0017-328	
SP	Organic	Hewlett Packard GC 5890 with ECD + ECD detectors and 7673A autosampler	2750A18077	GC			
SP	Organic	Hewlett Packard GC 5890 with ECD + ECD detectors and 7673A autosampler	2413A11796	GC			
SP	Organic	Hewlett Packard GC 5890 with FID + NPD detectors and 7673A autosampler	2415A01377	GC	204	26-91-0042	
SP	Organic	Hewlett Packard GC 5890 with NPD + FPD detectors and 7673A autosampler	2916A22297	GC	201		
SP	Organic	Hewlett Packard GC 5890 with PID + FID detectors and LSC2000/ALS2032 autosampler	2916A22291	GC	207	89-135	
SP	Organic	Hewlett Packard GC 5890 with PID + FID detectors and LSC2000/ALS2032 autosampler	323A42727	GC	213	26-95-0070-363	
SP	Organic	Hewlett Packard GC 5890 with ELCD + PID detectors and LSC2000/ALS2016 autosampler	2541A06581	GC	209	26-95-0018-342	
SP	Organic	Hewlett Packard GC 5890 with PID + FID detectors and LSC2000/ALS2016 autosampler	2541A06582	GC	203	26-95-0033-332	1

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FGL Environmental Instrument and Equipment List

I.AB	DIVISION	INSTRUMENT DESCRIPTION	SERIAL NUMBER	INST. TYPE	INST. ID	INV. CONTROL	REMARKS
SP	Organic	Hewlett Packard GC 5890 with 2 ELCS/PID detectors and LSC2000/ALS2016 autosampler	2950A28076	GC	211	26-95-0063-356	
SP	Organic	Hewlett Packard GC 5890 with 2 ELCS/PID detectors and LSC2000/ALS2016 autosampler	3115A35121	GC	212		
SP	Organic	Hewlett Packard GC 5890II with FID and and LSC2000/ALS2016 autosampler	3033A30381	GC		26-95-0038-334-1	
SP	Organic	Hewlett Packard GC 5890 with ELCD + ECD detectors and 7673 autosampler		GC			
SP	Organic	Hewlett Packard HPLC 1090 with UV and Fluorescence detector and post column derivatization		HPLC	202	89-126	
SP	Organic	Hewlett Packard HPLC 1090 with diode array and fluorescence detectors	2623A00932	HPLC	201		
SP	Organic	Hitachi HPLC L-6200A with L-450 Diode Array and Shimadzu RF-551 Fluorescence detector	0312-054	HPLC	203	26-92-0018	
SP	Organic	Foxboro Miran 1FF Infra-red Spectrophotometer	2539	IR	201	26-93-0041	
SP	Organic	Horiba OCMA-22O Infra-red Spectrophotometer	202021	IR	202		
SP	Organic	MCI TOX 10	43C30238	тох	201	88-099	
SP	Organic	ASTRO 2001	2390072501	тос	201	90-190	
SP	Organic	Mettler PC 440 Analytical Balance	814108	SCALE	206		
SP	Inorganic	Fisons Plasma Quad 2 ICP/MS	970	ICP/MS	201	26-93-00348	
SP	Inorganie	Hewlett Packard 4500 ICP/MS	3528J00253	ICP/MS	202	26-95-0057-350	
SP	Inorganic	ARL 3410 ICP with model 101 autosampler	1324	ICP	201	90-178	
SP	Inorganic	Thermo-Jerrell Ash Atomscan 16 ICP	1871	ICP	202	26-95-0001-323	
SP	Inorganic	Thermo-Jerrell Ash 16E Trace 61E	338490	ICP/TR	201	26-95-0048-344	
SP	Inorganic	Thermo-Jerrell Ash, JoE Purge 616	8290	ICP/PR	201		

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FGL Environmental Instrument and Equipment List

LAB	DIVISION	INSTRUMENT DESCRIPTION	SERIAL NUMBER	INST. TYPE	INST. ID	INV. CONTROL	REMARKS
SP	Inorganic	Perkin-Elmer 5100Z GFAA with AS-60 autosampler	136513	GFAA	201	89-134	
SP	Inorganic	Perkin-Elmer 5100Z GFAA with AS-60 autosampler	137543	GFAA	202	26-95-0061-354	
SP	Inorganic	Perkin-Elmer 5000 FAA	123286	FAA	201	82-051	
SP	Inorganic	LEEMAN PS200 Automated Mercury Analyzer	1156/110/00012	CVAA	201	26-96-0028-411	
SP	Inorganic	Dionex 300 Ion Chromatograph and Spectrophysics AS3500 autosampler	921104	IC	201	26-92-0014	
SP	Inorganic	Dionex 100 Ion Chromatograph and Spectrophysics AS3500 autosampler	94110001	IC	202	26-95-002-324	
SP	Inorganic	Technicon AA2 Flow injection autoanalyzer	FGL-09304REP	FIA	202		
SP	Inorganic	Technicon AA2 Flow injection autoanalyzer	FGL-08304REP	FIA	201		
SP	Inorganic	Perkin-Elmer Lambda 3 UV/VIS Spectrophotometer	46034	UV/VIS	201	81-043	
SP	Inorganic	Sequuia-Turner model 690 Nephelometer (Turbidimeters)	2426	TURBID	201	87-091	
SP	Inorganic	Orion model 860 Oxygen Meter	19090015	O2	201		
SP	Inorganic	Orion model 160 Conductivity Meter	09050045	EC	201	26-91-0027	
SP	Inorganic	Fisher Scientific Accumet pH Meter # 25	C0006984	PH	201		
SP	Inorganic	Setaflash	2010	FLASH	201		
sp	Inorganies	CEM MDS 2100 Microwave Digestor	Z4023	MICRO	201		
SP	Inorganic	Mettler AE 200 Analytical Balance	L19861	SCALE	202	26-93-0038	
SP	Inorganic	Mettler Toledo PB 303 Analytical Balance	1113242753	SCALE	201		
SP	Radio	Princeton Gamma Tech. Gamma Ray Spectroscope	2656	GS	201	26-94-0005	

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FGL Environmental Instrument and Equipment List

LAB	DIVISION	INSTRUMENT DESCRIPTION	SERIAL NUMBER	INST. TYPE	INST. ID	INV, CONTROL	REMARKS
SP	Radio	Oxford Tennelec Oasis System, Alpha Spectrometer	30104	AS	201-208	26-95-0012-327	
SP	Radio	Radom 5C-5, Gas Scintillation	13371270	SC	201	26-93-0018	
SP	Radio	Packard 2500TR, Liquid Scintillation	2500	LSC	201	26-93-197	
SP	Radio	NMC PCCIIT, Alpha Counter	79-2750-07	GP	201	26-79-999	
SP	Radio	NMC PCCIIT, Alpha Counter	81-2750-14	GP	202	26-81-045	
SP	Radio	NMC PCC11T, Alpha Counter	89-2750-1-17	GP	205	26-89-153-1	
SP	Radio	NMC PCC11T, Alpha Counter	89-2750-1-18	GP	206	26-89-153-2	
SP	Radio	NMC PCC11T, Alpha Counter	90-2755-1-03	GP	207	26-90-167	
SP	Radio	Tennelec LB 1000, Alpha/Beta Counter	577	GP	203	26-87-083	
SP	Radio	Tennelee LB 1000, Alpha/Beta Counter	518	GP	204	26-89-111	
SP	Radio	Tennelec LB 5100, Alpha/Beta Counter, with autosampler	505	GP	208	26-93-0033	
SP	Radio	Sartorius L 420 P Balance	37080053	SCALE	204	90-165	
SP	Bact	2 Incubator	N/A	N/A	N/A	N/A	
SP	Bact	2 Autoclaves	N/A	N/A	N/A	N/A	
SP	Field Service	10 Vehicles	N/A	N/A	N/A	N/A	
SP	Field Service	9 Isco Autosamplers	N/A	N/A	N/A	N/A	
SP	Computers	3 Microvax with PCSA fileserver using Foxpro for DOS	N/A	N/A	N/A	N/A	
SP	Computers	I WAN/LAN Xyplex Chassis	N/A	N/A	N/A	N/A	1

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FGL Environmental Instrument and Equipment List

LAB	DIVISION	INSTRUMENT DESCRIPTION	SERIAL NUMBER	INST. TYPE	INST. ID	INV. CONTROL	REMARKS
SP	Computers	1 BBS by Galacticomm	N/A	N/A	N/A	N/A	
SP	Computers	80 DOS Based Computers	N/A	N/A	N/A	N/A	
S1K	Organic	Hewlett Packard GC/MS 5890II/5972 with LSC2000/ALS2016 autosampler	2921A22867	GC/MS	301		
STK	Organic	Hewlett Packard GC 5890II with ECD + ECD detectors and 7673A autosampler	3223A43094	GC	302	46-93-0029-126	
STK	Organic	Hewlett Packard GC 5890 with FID + FID detectors and 7673A autosampler	2463A72072	GC	304		
STK	Organic	Hewlett Packard GC 5890 with PID + FID detectors and LSC2000/ALS2016 autosampler	3336A53365	GC	303	46-94-0033	
STK	Organic	Hewlett Packard GC 5890 with PID + ELCD detectors and LSC2000/ALS2016 autosampler	2750A15950	GC	301		
STK	Organic	Horiba, OCMA 220, Fixed IR	308023	IR	301	46-93-0051	
STK	Inorganic	Thermo-Jarrell Ash Atomscan 25	2508	1CP	301	46-93-0029	
STK	Inorganic	Hitachi 8270 Graphite Furnace AA with Zeeman Background Correction	0527-026	GFAA	302	46-94-0057	
STK	Inorganic	Perkin-Eliner 5100Z Graphite Furnace AA with Zeeman background correction and Flame AA	137931	GFAA	301	46-95-0041-131	
STK	Inorganic	Perkin-Eliner 5000 Flame AA set up for inercury cold vapor	117350		N/A		
STK	Inorganic	CEM MDS 2100 microwave digestor	7-4101	MICRO	301	46-93-0014	
STK	Inorganic	Dionex DX-300 Ion Chromatograph and Spectraphysics AS3500 autosampler	930309/013-0176	IC	301	46-93-0015	
STK	Inorganic	Technicaon AA2 Flow injection autoanalyzer	171-A017- 03/GC9683/GC12683	FIA	301		
STK	Inorganic	Orion model 860 Oxygen Meter	41555015	O2	301		

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FGL Environmental Instrument and Equipment List

LAB	DIVISION	INSTRUMENT DESCRIPTION	SERIAL NUMBER	INST. TYPE	INST. ID	INV. CONTROL	REMARKS
STK	Inorganic	Orion model 160 Conductivity Meter	44524019	EC	301	46-95-0003	
STK	Inorganic	Fisher Scientific Accumet pH Meter # 25	C0017099	PH	301		
STK	Inorganic	Sequoia-Turner 690 UV/VIS spectrophotometer	B011224	UV/VIS	301		
STK	Inorganic	Turner 40 Nephelometer	01151	TURBID	301		
STK	Bact	Incubator	N/A	N/A	N/A	N/A	
STK	Bact	Autoclave	N/A	N/A	N/A	N/A	
STK	Field Services	3 Vehicles	N/A	N/A	N/A	N/A	
STK	Field Services	3 Isco Autosamplers	N/A	N/A	N/A	N/A	
STK	Computers	1 Microvax with PCSA pathworks fileserver using Foxpro for DOS	N/A	N/A	N/A	N/A	
STK	Computers	1 Xyplex bridge router	N/A	N/A	N/A	N/A	
STK	Computers	20 DOS based computers	N/A	N/A	N/A	N/A	

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Revision No: 1.0

Date: September 22, 1995

Document Control and Records Management -

This section is currently in the format of the actual SOP for Document control. Formal revision into the proper format used throughout the QA Manual will be forthcoming. Also, this currently only covers the document control portion of records management. Total records management will be addressed in the next revision.

Scope and Application:

This procedure outlines the overall document control for the laboratory. This includes outlines for determining if the document needs control and how to remove documents and turnover to archive. These methods will prevent document users from referencing outdated documents.

Equipment:

computers, files, binders etc.

Procedure:

The important decision that needs to be made for all documents is whether or not the document needs to be controlled. The following are a few criteria for deciding whether the document needs to be controlled

- 1) Does the document need to be retained for proof of analysis or procedures used?
- 2) Will the document be revised over time?
- 3) Will more than one copy of the document be in use?

The following is a list of documents known to require document control:

Standard Operating Procedures Quality Assurance Manual Statement of Qualifications Fee Schedule

Document control:

Beginning control of a document:

- 1) Fill out the heading information on the Document Control Distribution Record Form.
- 2) Record the control number, who the document was issued to, the revision issued, and the date issued.
- 3) Stamp the front of the document with the controlled copy stamp and fill in the control number.

Continuing control of a document:

- 1) Upon the next revision of the document refer to the Document Control Distribution Record to determine who needs the new revision.
- 2) Stamp the front of the revised document with the controlled copy stamp and fill in the same control number if issued to the same recipient.
- 3) Issue the new controlled revision to each recipient of a controlled copy with a letter and return envelope requesting they return the outdated document.
- 4) The letter should also inquire if they need to remain on the list for receiving a controlled copy of the document.
- 5) Upon return of the controlled copy the Document Control Distribution Record should be updated recording that the document was returned. Also, document if the recipient no longer needs to receive a controlled copy.
- 6) One copy of the outdated document should be stamped with the superceded date stamp, dated and filed.
- 7) All other copies of the document should be destroyed after updating the Document Control Distribution Record.

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If a controlled document is released without document control it MUST be stamped INFORMATION COPY ONLY

Calculations:

N/A

Record Keeping and Storage:

There are two sets of files required for proper document control. Both sets of files must be strored in a central location under the supervision of the person responsible for upkeep. They are as follows:

- 1) Files for the Document Control Distribution Record Forms. A file for each department needs to be maintained for record storage.
- 2) Files for the superceded documents. A corresponding file must be created to store the superceded revisions of the document.

A list of files to be created can be easily obtained for the known documents requiring control. It is created as follows:

- 1) SOP's: All LIMS subdirectories under M:\FGLDOCS\SOP will require an individual file (which usually reflects the department).
- 2) QA Manual
- 3) Statement of Qualifications
- 4) Fee Schedule

Further documents to be controlled will be in future revisions of this document.

References:

N/A

Quality Assurance Manual - Amendment

Amendment Date: 8-20-96

Sections Amended: 2, 3, 4, 5, 6, 7, 9, 10, 11, 12, Appendix A

Reviewed by: Yee Ren, M.S.

Data Validator

Concurred by:

Kurt Wilkinson, B.S.

Quality Assurance Director

19/96

| Date | S/21/96 |
| Date | Date | S/31/96 |
| Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | Date | D Approved by: Darrell H. Nelson, B.S.

Laboratory Director

This amendment should be inserted at the back of the quality assurance manual. The next formal revision (scheduled for April, 1997) will incorporate these amendments into their respective sections of the manual.

APPENDIX F BLANK, DUPLICATE, AND SPIKE SAMPLE ANALYTICAL REPORTS

April 4, 1997

LAB No: SP 702024-1

Bermite Division of Whittaker 22116 W. Soledad Canyon Road RE: Organic Analysis Matrix: Monitoring Well

Saugus, CA 91350

Project/Site: MW5-1A Quarterly Sampling Area 317

Sample Description: MW5/B/34/1A Sampled : March 19, 1997 Sampled by : Abdun-Nur/Bricker Received : March 19, 1997

Preservatives: H2SO4 pH < 2 Analyzed : March 20, 1997

QA/QC ID# : SP 97032000A A

TOTAL ORGANIC CARBON

CONSTITUENT	EPA METHOD	UNITS	SAMPLE DLR	SAMPLE RESULTS	LAB DLR	BLANK RESULTS
TOC	415.1	mg/L	0.5	ND	0.5	ND

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.)
mg/L = Milligrams Per Liter (ppm)
ND = Not Detected at or above the DLR.

• = DLR adjusted because of dilutions, concentrations, or limited sample.

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

Kelly A. Dunnahoo, B.S. Organic Laboratory Manager

KAD:cea

Field Office Visalia, CA TEL: 209/734-9473 FAX: 209/734-8435

April 4, 1997

LAB No: SP 702024-2

Bermite Division of Whittaker 22116 W. Soledad Canyon Road RE: Organic Analysis Matrix: Monitoring Well

Saugus, CA 91350

Project/Site: MW5-1A Quarterly Sampling Area 317

Sample Description: MW5/C/34/1A
Sampled by: Abdun-Nur/Bricker

Sampled : March 19, 1997 Received : March 19, 1997

Container : Amber Glass TFE-Cap

Extracted: N/A

Preservatives:

Analyzed : March 25, 1997 QA/QC ID# : SP 97032500A A

TOTAL ORGANIC HALOGENS

CONSTITUENT	EPA METHOD	UNITS	SAMPLE DLR	SAMPLE RESULTS	LAB DLR	BLANK RESULTS
TOX	9020	ug/L	5	ND	5	ND

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.)
ug/L = Micrograms Per Liter (ppb)
ND = Not Detected at or above the DLR.

• = DLR adjusted because of dilutions, concentrations, or limited sample.

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

Kélly A. Dunnahoo, B.S. Organic Laboratory Manager

April 4, 1997

LAB No: SP 702024-3

Bermite Division of Whittaker 22116 W. Soledad Canyon Road

RE: Organic Analysis
Matrix: Monitoring Well

Saugus, CA 91350

Project/Site: MW5-1A Quarterly Sampling Area 317

Sample Description: MW5/0/34/1A Sampled by : Abdun-Nur/Bricker

Sampled : March 19, 1997 Received : March 19, 1997

Container : VOA

Extracted: N/A

Preservatives:

Analyzed : March 21, 1997 QA/QC ID# : SP 97032101H A

EPA METHOD 624

CONSTITUENT	SAMPLE	SAMPLE	LAB BLANK
	DLR	RESULTS	DLR RESULTS
	ug/L	ug/L	ug/L ug/L
Trichloroethylene	0.5	ND	0.5 ND
SURROGATES	SAI	MPLE	LAB BLANK
	AR	% REC.	AR % REC.
1,2-Dichloroethane-d4	40-140	120	40-140 125
Toluene-d8	64-139	103	64-139 102
BFB	50-149	109	50-149 108

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.)

ug/L = Micrograms Per Liter (ppb)

ND = Not Detected at or above the DLR. AR = Acceptable Range

• = DLR adjusted because of dilutions, concentrations, or limited sample.

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

Kelly A. Dunnahoo, B.S. Organic Laboratory Manager



Analytical Chemists

April 4, 1997

ORGANIC Quality Assurance Report for sample: 702024

Bermite Division of Whittaker 22116 W. Soledad Canyon Road Saugus , CA 91350

	BATCH	EPA			CALIB	RATION	QA/QC					METHOD	QA/QC			
Constituent	ID	Method	Units	Туре	Conc.	% REC	AR	NOTE	Type	Conc.	% REC	% REC	AR	% DIF	MAV	NOTE
1,2-Dichloroethane-d4	1H12A	624	ug/L	LCS	10.0	116	40-140		BS	10.0	119	121	40-140	1.0	30.0	
Toluene-d8	1H12A	624	ug/L	LCS	10.0	105	64-139		BS	10.0	104	104	64-139	0.0	30.0	
BFB	1H12A	624	ug/L	LCS	10.0	111	50-149		BS	10.0	110	110	50-149	0.5	30.0	
Trichloroethylene	1H12A	624	ug/L	LCS	10.0	106	37-151		BS	10.0	108	108	37-151	0.2	23.0	
TOC	0A 2A	415.1	mg/L	LCS	55.0	89.1	75-125		MS	50.0	92.0	98.0	75-125	6.3	20.0	
тох	5A 2A	TOX	ug/L	CCA	10.0	91.0	74-116		MS	100	91.0	101	75-125	10.4	20.0	

FGL ID = 19970320 N/A => Not Applicable NOTE => See note indicated below.

FGL ENVIRONMENTAL, INC.

KAD:cea

Kurt Wilkinson, B.S., QA Director

April 4, 1997

LAB No: SP 702026-1

Bermite Division of Whittaker 22116 W. Soledad Canyon Road

RE: Organic Analysis
Matrix: Monitoring Well

Saugus , CA 91350

Project/Site: MW6-1A Quarterly Sampling Area 317

Sample Description: MW6/B/34/1A Sampled : March 19, 1997 Sampled by : Abdun-Nur/Bricker Received : March 19, 1997

Preservatives: H2SO4 pH < 2 Analyzed : March 20, 1997 QA/QC ID# : SP 97032000A A

TOTAL ORGANIC CARBON

CONSTITUENT	EPA METHOD	UNITS	SAMPLE DLR	SAMPLE RESULTS	LAB DLR	BLANK RESULTS
TOC	415.1	mg/L	0.5	ND	0.5	ND

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.)
mg/L = Milligrams Per Liter (ppm)
ND = Not Detected at or above the DLR.

♦ = DLR adjusted because of dilutions, concentrations, or limited sample.

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

Kelly A. Dunnahoo, B.S. Organic Laboratory Manager

April 4, 1997

LAB No: SP 702026-2

Bermite Division of Whittaker 22116 W. Soledad Canyon Road RE: Organic Analysis
Matrix: Monitoring Well

Saugus, CA 91350

Project/Site: MW6-1A Quarterly Sampling Area 317

Sample Description: MW6/C/34/1A Sampled by : Abdun-Nur/Bricker

Sampled : March 19, 1997 Received : March 19, 1997

Container : Amber Glass TFE-Cap

Extracted : N/A

Preservatives:

Analyzed: March 25, 1997

QA/QC ID# : SP 97032500A A

TOTAL ORGANIC HALOGENS

CONSTITUENT	EPA METHOD	UNITS	SAMPLE DLR	SAMPLE RESULTS	LAB DLR	BLANK RESULTS
тох	9020	ug/L	5	ND	5	ND

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.)
ug/L = Micrograms Per Liter (ppb)

ND = Not Detected at or above the DLR.

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

Kelly A. Dunnahoo, B.S. Organic Laboratory Manager

^{• =} DLR adjusted because of dilutions, concentrations, or limited sample.

April 4, 1997

LAB No: SP 702026-3

Bermite Division of Whittaker 22116 W. Soledad Canyon Road

RE: Organic Analysis Matrix: Monitoring Well

Saugus , CA 91350

Project/Site: MW6-1A Quarterly Sampling Area 317

Sample Description: MW6/0/34/1A Sampled by : Abdun-Nur/Bricker

Sampled : March 19, 1997 Received : March 19, 1997

Container : VOA

Extracted: N/A

Preservatives:

Analyzed : March 21, 1997 QA/QC ID# : SP 97032101H A

EPA METHOD 624

CONSTITUENT	SAMPLE	SAMPLE	LAB BLANK
	DLR	RESULTS	DLR RESULTS
	ug/L	ug/L	ug/L ug/L
Trichloroethylene	0.5	ND	0.5 ND
SURROGATES	SAI	MPLE	LAB BLANK
	AR	% REC.	AR % REC.
1,2-Dichloroethane-d4	40-140	120	40-140 125
Toluene-d8	64-139	103	64-139 102
BFB	50-149	107	50-149 108

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.)

ug/L = Micrograms Per Liter (ppb)

ND = Not Detected at or above the DLR. AR = Acceptable Range

• = DLR adjusted because of dilutions, concentrations, or limited sample.

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

Kelly A. Dunnahoo, B.S. Organic Laboratory Manager



Analytical Chemists

April 4, 1997

ORGANIC Quality Assurance Report for sample: 702026

Bermite Division of Whittaker 22116 W. Soledad Canyon Road Saugus , CA 91350

	ВАТСН	EPA			CALIE	RATION	QA/QC					METHOD	QA/QC			
Constituent	ID	Method	Units	Туре	Conc.	% REC	AR	NOTE	Type	Conc.	% REC	% REC	AR	% DIF	MAV	NOTE
1,2-Dichloroethane-d4	1H12A	624	ug/L	LCS	10.0	116	40-140		BS	10.0	119	121	40-140	1.0	30.0	
Toluene-d8	1H12A	624	ug/L	LCS	10.0	105	64-139		BS	10.0	104	104	64-139	0.0	30.0	
BFB	1H12A	624	ug/L	LCS	10.0	111	50-149		BS	10.0	110	110	50-149	0.5	30.0	
Trichloroethylene	1H12A	624 .	ùg/L	LCS	10.0	106	37-151	, .	BS	10.0	108	108	37-151	0.2	23.0	
тос	OA 2A	415.1	mg/L	LCS	55.0	89.1	75-125		MS	50.0	92.0	98.0	75-125	6.3	20.0	
TOX	5A 2A	TOX	ug/L	CCA	10.0	91.0	74-116		MS	100	91.0	101	75-125	10.4	20.0	

FGL ID = 19970320 N/A => Not Applicable NOTE => See note indicated below.

FGL ENVIRONMENTAL, INC.

KAD:cea

Kurt Wilkinson, B.S., QA Director

APPENDIX G ANALYTICAL REPORTS FOR GROUND WATER MONITORING PARAMETERS

April 3, 1997

Bermite Division of Whittaker 22116 W. Soledad Canyon Road Saugus , CA 91350 LAB No: SP 702021-1

RE: Inorganic Analysisy

ECEIVE APR 1 4 1997

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Sample Site: MW1 Quarterly Sampling Area 317

Description: MW1/A/34

Sampled by : Abdun-Nur/Bricker Type of Sample: Monitoring Well Sampled: March 19, 1997 Received: March 19, 1997 Completed: March 21, 1997

QA/QC ID# : 70202101-

Analytical Results

CONSTITUENT	METHOD	UNITS	DLR	RESULTS
Conductivity	2510B	umhos/cm2	1	770
pH	4500-H B	units		7.8

DLR = Detection Limit for Reporting Purposes. ND = Not Detected at or above the DLR.

ug/L = Micrograms Per Liter (ppb) mg/L = Milligrams Per Liter (ppm) mg/kg = Milligrams Per Kilogram

• = DLR adjusted because of dilutions, concentrations, or limited sample.

Preservatives: (1) Cool 4°C Containers: (a) Plastic

If you have any questions, please call.

FGL ENVIRONMENTAL

Kurt Wilkinson, B.S.

QA Director

April 3, 1997

LAB No: SP 702021-2

Bermite Division of Whittaker 22116 W. Soledad Canyon Road RE: Organic Analysis Matrix: Monitoring Well

Saugus , CA 91350

Project/Site: MW1 Quarterly Sampling Area 317

Sample Description: MW1/B/34 Sampled : March 19, 1997 Sampled by : Abdun-Nur/Bricker Received : March 19, 1997

Container: Amber Glass TFE-Cap Extracted: N/A

Preservatives: H2SO4 pH < 2

Analyzed: March 20, 1997

QA/QC ID# : SP 97032000A A

TOTAL ORGANIC CARBON

CONSTITUENT	EPA METHOD	UNITS	SAMPLE DLR	SAMPLE RESULTS	LAB DLR	BLANK RESULTS
TOC	415.1	mg/L	0.5	ND	0.5	ND

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.)
mg/L = Milligrams Per Liter (ppm)
ND = Not Detected at or above the DLR.

• = DLR adjusted because of dilutions, concentrations, or limited sample.

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

Kelly A. Dunnahoo, B.S. Organic Laboratory Manager

April 3, 1997

LAB No: SP 702021-3

Bermite Division of Whittaker 22116 W. Soledad Canyon Road RE: Organic Analysis Matrix: Monitoring Well

Saugus, CA 91350

Project/Site: MW1 Quarterly Sampling Area 317

Sample Description: MW1/C/34
Sampled by: Abdun-Nur/Bricker

Sampled : March 19, 1997 Received : March 19, 1997

Container : Amber Glass TFE-Cap

Extracted: N/A

Preservatives:

Analyzed : March 25, 1997 QA/QC ID# : SP 97032500A A

TOTAL ORGANIC HALOGENS

CONSTITUENT	EPA METHOD	UNITS	SAMPLE DLR	SAMPLE RESULTS	LAB DLR	BLANK RESULTS
тох	9020	ug/L	5	ND	5	ND

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.)
ug/L = Micrograms Per Liter (ppb)

ND = Not Detected at or above the DLR.

• = DLR adjusted because of dilutions, concentrations, or limited sample.

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

Kelly A. Dunnahoo, B.S. Organic Laboratory Manager

KAD:cea

Field Office Visalia, CA TEL: 209/734-9473

April 3, 1997

LAB No: SP 702021-4

Bermite Division of Whittaker 22116 W. Soledad Canvon Road

RE: Inorganic Analysis

Saugus, CA 91350

Sample Site: MW1 Quarterly Sampling Area 317

Description: MW1/H/34

Sampled: March 19, 1997

Sampled by : Abdun-Nur/Bricker Type of Sample: Monitoring Well Received: March 19, 1997

ell Completed

Completed: March 27, 1997

QA/QC ID# : 70202104-

Analytical Results

CONSTITUENT	METHOD	UNITS	DLR	RESULTS
Chloride	4110B	mg/L	2.0*	170
Sulfate	4110B	mg/L		10

DLR = Detection Limit for Reporting Purposes. ND = Not Detected at or above the DLR.

ug/L = Micrograms Per Liter (ppb) mg/L = Milligrams Per Liter (ppm) mg/kg = Milligrams Per Kilogram

• = DLR adjusted because of dilutions, concentrations, or limited sample.

Preservatives: (1) Cool 4°C Containers: (a) Plastic

If you have any questions, please call.

FGL ENVIRONMENTAL

Kurt Wilkinson, B.S.

QA Director

April 3, 1997

LAB No: SP 702021-5

Extracted: N/A

Bermite Division of Whittaker 22116 W. Soledad Canyon Road

RE: Organic Analysis Matrix: Monitoring Well

Saugus , CA 91350

Project/Site: MW1 Quarterly Sampling Area 317

Sample Description: MW1/0/34 Sampled : March 19, 1997 Sampled by : Abdun-Nur/Bricker Received : March 19, 1997

Container : VOA

Preservatives:

Analyzed: March 21, 1997
QA/QC ID#: SP 97032101H A

EPA METHOD 624

CONSTITUENT	SAMPLE	SAMPLE	LAB BLANK
	DLR	RESULTS	DLR RESULTS
	ug/L	ug/L	ug/L ug/L
Trichloroethylene	0.5	ND	0.5 ND
SURROGATES	SAI	MPLE	LAB BLANK
	AR	% REC.	AR % REC.
1,2-Dichloroethane-d4	40-140	122	40-140 125
Toluene-d8	64-139	103	64-139 102
BFB	50-149	108	50-149 108

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.)

ug/L = Micrograms Per Liter (ppb)

ND = Not Detected at or above the DLR. AR = Acceptable Range

• = DLR adjusted because of dilutions, concentrations, or limited sample.

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

Kelly A. Dunnahoo, B.S. Organic Laboratory Manager

KAD:cea

Field Office Visalia, CA TEL: 209/734-9473

April 3, 1997

LAB No: SP 702021-6

Bermite Division of Whittaker

22116 W. Soledad Canyon Road

Saugus, CA 91350

RE: Inorganic Analysis

Sample Site: MW1 Quarterly Sampling Area 317

Description: MW1/R/34

: March 19, 1997 Sampled

Sampled by : Abdun-Nur/Bricker Type of Sample: Monitoring Well Received: March 19, 1997 Completed: March 31, 1997

QA/QC ID# : 70202106-

Analytical Results

CONSTITUENT	METHOD	UNITS	DLR	RESULTS	MCL
Iron Manganese Sodium	200.7 200.8 200.7	ug/L ug/L mg/L	50 0.5 1	100 3.2 53	300 50

DLR = Detection Limit for Reporting Purposes. ND = Not Detected at or above the DLR.

ug/L = Micrograms Per Liter (ppb) mg/L = Milligrams Per Liter (ppm) mg/kg = Milligrams Per Kilogram

◆ = DLR adjusted because of dilutions, concentrations, or limited sample. Preservatives: (1) Cool 4°C (2) HNO3 pH < 2 Containers: (a) Plastic

If you have any questions, please call.

FGL ENVIRONMENTAL

Kurt Wilkinson, B.S.

QA Director



Analytical Chemists

April 3, 1997

INORGANIC Quality Assurance Report for sample: 702021

Bermite Division of Whittaker 22116 W. Soledad Canyon Road Saugus , CA 91350

	BATCH			BL	LANK QA/QC			CALIE	RATION	QA/QC					METHOD	QA/QC			
Constituent	ID	Method	Units	DLR	Result	NOTE	Туре	Conc.	% REC	AR	NOTE	Type	Conc.	% REC	% REC	AR	% DIF	MAV	NOTE
Iron	2A 2A	200.7	ug/L	50	ND		ccv	5000	100	90-110		MS	500	99.0	99.3	75-125	0.3	20.0	
Manganese	12E 2A	200.8	ug/L-mg/L	0.50	ND		CCV	100	98.4	90-110		MS	80.0	94.0	90.2	75-125	3.9	20.0	=
Sodium	2A 2A	200.7	mg/L	1.0	ND		CCV	25.0	97.6	90-110		MS	2.50	70.0	70.8	75-125	0.0	20,0	405
Chloride ·	8A · 2D	4110B	mg/L '	1.0	ND	•	CCV	50.0	103	90-110	•	MS	400	(32.4)	5.4	85-111	43.7	4.1	410
Conductivity	2A 2A	2510B	umhos/c2	1.0	ND		CCV	1000	98.0	75-125		Dup	1160	N/A	N/A	N/A	0.4	0.8	
pH	GA 2A	4500-H B	units		N/A		CCV	8.00	99.8	90-110		Dup	7.75	N/A	N/A	N/A	0.5	1.4	
Sulfate	8A 2C	4110B	mg/L	1.0	ND		CCV	100	92.6	90-110		MS	400	96.8	90.9	80-120	6.1	20.0	

FGL ID = 19970319 ND => Not Detected at ar above DLR. DLR => Detection Limit for Reporting purposes. N/A => Not Applicable NOTE => See note indicated below.

Notes:

405 Matrix Spike (MS) not within the Acceptance Range (AR) and/or Relative Percent Difference (RPD) not within the Maximum Allowable Value (MAV) because of low concentrations of the spike or sample. Batch qualified based on the LCS, CCV or ICV recovery.

410 Matrix Spike (MS) not within Acceptance Range (AR) and/or Relative Percent Difference (RPD) not within Maximum Allowable Value (MAV). Batch qualified based on the LCS, CCV, or ICV recovery.

FGL ENVIRONMENTAL, INC.

KW:cea

Kurt Wilkinson, B.S., QA Director



Analytical Chemists

April 3, 1997

ORGANIC Quality Assurance Report for sample: 702021

Bermite Division of Whittaker 22116 W. Soledad Canyon Road Saugus , CA 91350

	BATCH	EPA			CALIE	RATION	QA/QC					METHOD	QA/QC			
Constituent	ID	Method	Units	Туре	Conc.	% REC	AR	NOTE	Туре	Conc.	% REC	% REC	AR	% DIF	MAV	NOTE
1,2-Dichloroethane-d4	1H12A	624	ug/L	LCS	10.0	116	40-140		BS	10.0	119	121	40-140	1.0	30.0	
Toluene-d8	1H12A	624	ug/L	LCS	10.0	105	64-139		BS	10.0	104	104	64-139	0.0	30.0	
BFB	1H12A	624	ug/L	LCS	10.0	111	50-149		BS	10.0	110	110	50-149	0.5	30.0	
Trichloroethylene .	1H12A	624	ug/L	LCS	10.0	106	37-151	•	BS	10.0	108	108	37-151	0.2	23.0	
TOC	0A 2A	415.1	mg/L	LCS	55.0	89.1	75-125		MS	50.0	92.0	98.0	75-125	6.3	20.0	
TOX	5A 2A	TOX	ug/L	CCV	10.0	91.0	74-116		MS	100	91.0	101	75-125	10.4	20.0	

FGL ID = 19970320 N/A => Not Applicable NOTE => See note indicated below.

FGL ENVIRONMENTAL, INC.

KAD:cea

Kurt Wilkinson, B.S., QA Director

April 3, 1997

LAB No: SP 702022-1

Bermite Division of Whittaker 22116 W. Soledad Canyon Road RE: Inorganic Analysis

Saugus , CA 91350

Sample Site: MW3 Quarterly Sampling Area 317

Description: MW3/A/34
Sampled by : Abdun-Num/Bricker

Sampled: March 19, 1997 Received: March 19, 1997

Sampled by : Abdun-Nur/Bricker Type of Sample: Monitoring Well

Completed: March 21, 1997

QA/QC ID# : 70202201-

Analytical Results

CONSTITUENT	METHOD	UNITS	DLR	RESULTS
Conductivity	2510B	umhos/cm2	1	600
pH	4500-H B	units		7.6

DLR = Detection Limit for Reporting Purposes. ND = Not Detected at or above the DLR. ug/L = Micrograms Per Liter (ppb) mg/L = Milligrams Per Liter (ppm) mg/kg = Milligrams Per Kilogram

♦ = DLR adjusted because of dilutions, concentrations, or limited sample.

Preservatives: (1) Cool 4°C (2) H2SO4 pH < 2 Containers: (a) Plastic

If you have any questions, please call.

FGL ENVIRONMENTAL

Kurt Wilkinson, B.S.

QA Director

April 3, 1997

LAB No: SP 702022-2

Bermite Division of Whittaker 22116 W. Soledad Canyon Road RE: Organic Analysis Matrix: Monitoring Well

Saugus, CA 91350

Project/Site: MW3 Quarterly Sampling Area 317

Sample Description: MW3/B/34 Sampled : March 19, 1997 Sampled by : Abdun-Nur/Bricker Received : March 19, 1997

Container: Amber Glass TFE-Cap Extracted: N/A

Preservatives: H2SO4 pH < 2 Analyzed : March 20, 1997 OA/OC ID# : SP 97032000A A

TOTAL ORGANIC CARBON

CONSTITUENT	EPA METHOD	UNITS	SAMPLE DLR	SAMPLE RESULTS	LAB DLR	BLANK RESULTS
TOC	415.1	mg/L	0.5	ND	0.5	ND

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.)
mg/L = Milligrams Per Liter (ppm)
ND = Not Detected at or above the DLR.

• = DLR adjusted because of dilutions, concentrations, or limited sample.

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

Kelly A. Dunnahoo, B.S. Organic Laboratory Manager

April 3, 1997

LAB No: SP 702022-3

Bermite Division of Whittaker 22116 W. Soledad Canyon Road RE: Organic Analysis Matrix: Monitoring Well

Saugus , CA 91350

Project/Site: MW3 Quarterly Sampling Area 317

Sample Description: MW3/C/34
Sampled by: Abdun-Nur/Bricker
Container: Amber Glass TFE-Cap

Sampled : March 19, 1997 Received : March 19, 1997

Extracted : N/A

Preservatives:

Analyzed : March 25, 1997 QA/QC ID# : SP 97032500A A

TOTAL ORGANIC HALOGENS

CONSTITUENT	EPA METHOD	UNITS	SAMPLE DLR	SAMPLE RESULTS	LAB DLR	BLANK RESULTS
TOX	9020	ug/L	5	ND	5	ND

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.)

ug/L = Micrograms Per Liter (ppb)

ND = Not Detected at or above the DLR.

• = DLR adjusted because of dilutions, concentrations, or limited sample.

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

Kelly A. Dunnahoo, B.S. Organic Laboratory Manager

April 3, 1997

LAB No: SP 702022-4

Bermite Division of Whittaker 22116 W. Soledad Canyon Road

Saugus, CA 91350

RE: Inorganic Analysis

Sample Site: MW3 Quarterly Sampling Area 317

Description: MW3/H/34

Sampled : March 19, 1997 Received : March 19, 1997

Sampled by : Abdun-Nur/Bricker Type of Sample: Monitoring Well

Received: March 19, 1997 Completed: March 27, 1997

QA/QC ID# : 70202204-

Analytical Results

CONSTITUENT	METHOD	UNITS	DLR	RESULTS
Chloride	4110B	mg/L	1	29
Sulfate	4110B	mg/L	1	71

DLR = Detection Limit for Reporting Purposes. ND = Not Detected at or above the DLR.

ug/L = Micrograms Per Liter (ppb) mg/L = Milligrams Per Liter (ppm) mg/kg = Milligrams Per Kilogram

♦ = DLR adjusted because of dilutions, concentrations, or limited sample. Preservatives: (1) Cool 4°C (2) H2SO4 pH < 2 Containers: (a) Plastic</p>

If you have any questions, please call.

FGL ENVIRONMENTAL

Kurt Wilkinson, B.S.

OA Director

April 3, 1997

LAB No: SP 702022-5

Bermite Division of Whittaker 22116 W. Soledad Canyon Road RE: Organic Analysis Matrix: Monitoring Well

Saugus, CA 91350

Project/Site: MW3 Quarterly Sampling Area 317

Sample Description: MW3/0/34 Sampled : March 19, 1997 Sampled by : Abdun-Nur/Bricker Received : March 19, 1997

Container: VOA Extracted: N/A

Preservatives: Analyzed: March 21, 1997 QA/QC ID#: SP 97032101H A

EPA METHOD 624

CONSTITUENT	SAMPLE	SAMPLE	LAB BLANK
	DLR	RESULTS	DLR RESULTS
	ug/L	ug/L	ug/L ug/L
Trichloroethylene	0.5	ND	0.5 ND
SURROGATES	SAI	MPLE	LAB BLANK
	AR	% REC.	AR % REC.
1,2-Dichloroethane-d4	40-140	122	40-140 125
Toluene-d8	64-139	103	64-139 102
BFB	50-149	116	50-149 108

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.)

ug/L = Micrograms Per Liter (ppb)

ND = Not Detected at or above the DLR. AR = Acceptable Range

• = DLR adjusted because of dilutions, concentrations, or limited sample.

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

. Kelly A. Dunnahoo, B.S. Organic Laboratory Manager

April 3, 1997

LAB No: SP 702022-6

Bermite Division of Whittaker 22116 W. Soledad Canyon Road

Saugus, CA 91350

RE: Inorganic Analysis

Sample Site: MW3 Quarterly Sampling Area 317

Description: MW3/R/34

Sampled: March 19, 1997

Sampled by : Abdun-Nur/Bricker Type of Sample: Monitoring Well Received: March 19, 1997 Completed: March 31, 1997

QA/QC ID# : 70202206-

Analytical Results

CONSTITUENT	METHOD	UNITS	DLR	RESULTS	MCL
Iron Manganese Sodium	200.7 200.8 200.7	ug/L ug/L mg/L	50 0.5 1	ND ND 60	300 50

DLR = Detection Limit for Reporting Purposes. ND = Not Detected at or above the DLR.

ug/L = Micrograms Per Liter (ppb) mg/L = Milligrams Per Liter (ppm) mg/kg = Milligrams Per Kilogram

◆ = DLR adjusted because of dilutions, concentrations, or limited sample. Preservatives: (1) Cool 4°C (2) HNO3 pH < 2 Containers: (a) Plastic

If you have any questions, please call.

FGL ENVIRONMENTAL

Kurt Wilkinson, B.S.

QA Director



Analytical Chemists

April 3, 1997

INORGANIC Quality Assurance Report for sample: 702022

Bermite Division of Whittaker 22116 W. Soledad Canyon Road Saugus , CA 91350

Constituent	BATCH ID	Method	Units	BL DLR	ANK QA/QC Result	NOTE	Туре	CALIE Conc.	RATION % REC	QA/QC AR	NOTE	Туре	Conc.	% REC		QA/QC AR	% DIF	MAV	NOTE
Iron	2A 2A	200.7	ug/L	50	ND		CCV	5000	100	90-110		MS	500	99.0	99.3	75-125	0.3	20.0	
Manganese	12E 2A	200.8	ug/L-mg/L	0.50	ND		CCV	100	98.4	90-110		MS	80.0	94.0	90.2	75-125	3.9	20.0	
Sodium	ZA ZA	200.7	mg/L	1.0	ND		CCV	25.0	97.6	90-110		MS	2.50	70.0	70.8	75-125	0.0	20.0	405
Chloride	8A 2C	4110B	mg/L	1.0	NTD .	•	CCV .	50.0	91.9	90-110		MS	400	108	99.7	85-111	5.7	4.1	410
Conductivity	2A 2A	2510B	umhos/c2	1.0	ND		CCV	1000	98.0	75-125		Dup	1160	N/A	N/A	N/A	0.4	0.8	
pH	OA 2A	4500-н в	units		N/A		CCV	8.00	99.8	90-110		Dup	7.75	N/A	N/A	N/A	0.5	1.4	
Sulfate	8A 2C	4110B	mg/L	1.0	ND		ccv	100	92.6	90-110		MS	400	96.8	90.9	80-120	6.1	20.0	

FGL ID = 19970319 ND => Not Detected at ar above DLR. DLR => Detection Limit for Reporting purposes. N/A => Not Applicable NOTE => See note indicated below.

Notes:

405 Matrix Spike (MS) not within the Acceptance Range (AR) and/or Relative Percent Difference (RPD) not within the Maximum Allowable Value (MAV) because of low concentrations of the spike or sample. Batch qualified based on the LCS, CCV or ICV recovery.

410 Matrix Spike (MS) not within Acceptance Range (AR) and/or Relative Percent Difference (RPD) not within Maximum Allowable Value (MAV). Batch qualified based on the LCS, CCV, or ICV recovery.

FGL ENVIRONMENTAL, INC.

KW:cea

Kurt Wilkinson, B.S., QA Director



Analytical Chemists

April 3, 1997

ORGANIC Quality Assurance Report for sample: 702022

Bermite Division of Whittaker 22116 W. Soledad Canyon Road Saugus , CA 91350

Constituent	BATCH ID	EPA Method	Units	Туре	CALIB Conc.	RATION % REC	QA/QC AR	NOTE	Туре	Conc.	% REC	METHOD % REC	QA/QC AR	% DIF	MAV	NOTE
1,2-Dichloroethane-d4	1H12A	624	ug/L	LCS	10.0	116	40-140		BS	10.0	119	121	40-140	1.0	30.0	
Toluene-d8	1H12A	624	ug/L	LCS	10.0	105	64-139		BS	10.0	104	104	64-139	0.0	30.0	
BFB	1H12A	624	ug/L	LCS	10.0	111	50-149		BS	10.0	110	110	50-149	0.5	30.0	
Trichloroethylene	"1H12A	624	ug/L	LCS	10.0	106	37-151		BS	. 10.0	108	108	37-151	0.2	23.0	` .
тос	0A 2A	415.1	mg/L	LCS	55.0	89.1	75-125		MS	50.0	92.0	98.0	75-125	6.3	20.0	
TOX	5A 2A	TOX	ug/L	CCA	10.0	91.0	74-116		MS	100	91.0	101	75-125	10.4	20.0	

FGL ID = 19970320 N/A => Not Applicable NOTE => See note indicated below.

KAD:cea

Kurt Wilkinson, B.S., QA Director

FGL ENVIRONMENTAL, INC.

April 3, 1997

LAB No: SP 702023-1

Bermite Division of Whittaker 22116 W. Soledad Canyon Road RE: Inorganic Analysis

Saugus, CA 91350

Sample Site: MW5 Quarterly Sampling Area 317

Description: MW5/A/34 Sampled by : Abdun-Nur/Bricker Type of Sample: Monitoring Well Sampled: March 19, 1997 Received: March 19, 1997

Completed: March 21, 1997

QA/QC ID# : 70202301-

Analytical Results

CONSTITUENT	METHOD	UNITS	DLR	RESULTS
Conductivity	2510B	umhos/cm2	1	530
pH	4500-H B	units		7.8

DLR = Detection Limit for Reporting Purposes. ND = Not Detected at or above the DLR.

ug/L = Micrograms Per Liter (ppb) mg/L = Milligrams Per Liter (ppm) mg/kg = Milligrams Per Kilogram

• = DLR adjusted because of dilutions, concentrations, or limited sample.

Preservatives: (1) Cool 4°C Containers: (a) Plastic

If you have any questions, please call.

FGL ENVIRONMENTAL

Kurt Wilkinson, B.S.

QA Director

April 3, 1997

LAB No: SP 702023-2

Bermite Division of Whittaker 22116 W. Soledad Canyon Road

RE: Organic Analysis Matrix: Monitoring Well

Saugus, CA 91350

Project/Site: MW5 Quarterly Sampling Area 317

Sample Description: MW5/B/34 Sampled : March 19, 1997 Sampled by : Abdun-Nur/Bricker Received : March 19, 1997

Preservatives: H2SO4 pH < 2

Analyzed: March 20, 1997

QA/QC ID# : SP 97032000A A

TOTAL ORGANIC CARBON

CONSTITUENT	EPA METHOD	UNITS	SAMPLE DLR	SAMPLE RESULTS	LAB DLR	BLANK RESULTS
ТОС	415.1	mg/L	0.5	ND	0.5	ND

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.)

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

Kelly A. Dunnahoo, B.S. Organic Laboratory Manager

April 3, 1997

LAB No: SP 702023-3

Bermite Division of Whittaker 22116 W. Soledad Canyon Road

RE: Organic Analysis Matrix: Monitoring Well

Saugus , CA 91350

Project/Site: MW5 Quarterly Sampling Area 317

Sample Description: MW5/C/34 Sampled : March 19, 1997 Sampled by : Abdun-Nur/Bricker Received : March 19, 1997

Preservatives:

Analyzed : March 25, 1997 QA/QC ID# : SP 97032500A A

TOTAL ORGANIC HALOGENS

CONSTITUENT	EPA METHOD	UNITS	SAMPLE DLR	SAMPLE RESULTS	LAB DLR	BLANK RESULTS
TOX	9020	ug/L	5	ND	5	ND

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.)

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

Kelly A. Dunnahoo, B.S. Organic Laboratory Manager

April 3, 1997

LAB No: SP 702023-4

Bermite Division of Whittaker 22116 W. Soledad Canyon Road

Saugus, CA 91350

RE: Inorganic Analysis

Sample Site: MW5 Quarterly Sampling Area 317

Description: MW5/H/34

Sampled: March 19, 1997

Sampled by : Abdun-Nur/Bricker Type of Sample: Monitoring Well

Received: March 19, 1997 Completed: March 27, 1997

QA/QC ID# : 70202304-

Analytical Results

CONSTITUENT	METHOD	UNITS	DLR	RESULTS
Chloride	4110B	mg/L	1	54
Sulfate	4110B	mg/L	1	30

DLR = Detection Limit for Reporting Purposes. ND = Not Detected at or above the DLR.

ug/L = Micrograms Per Liter (ppb) mg/L = Milligrams Per Liter (ppm) mg/kg = Milligrams Per Kilogram

• = DLR adjusted because of dilutions, concentrations, or limited sample.

Preservatives: (1) Cool 4°C Containers: (a) Plastic

If you have any questions, please call.

FGL ENVIRONMENTAL

Kurt Wilkinson, B.S.

QA Director

ANALYTICAL CHEMISTS

April 3, 1997

LAB No: SP 702023-5

Bermite Division of Whittaker 22116 W. Soledad Canyon Road

RE: Organic Analysis Matrix: Monitoring Well

Saugus , CA 91350

Project/Site: MW5 Quarterly Sampling Area 317

Sample Description: MW5/0/34 Sampled by : Abdun-Nur/Bricker

Sampled: March 19, 1997 Received: March 19, 1997

Container : VOA Preservatives:

Extracted: N/A

Analyzed : March 21, 1997 QA/QC ID# : SP 97032101H A

EPA METHOD 624

CONSTITUENT	SAMPLE	SAMPLE	LAB BLANK
	DLR	RESULTS	DLR RESULTS
	ug/L	ug/L	ug/L ug/L
Trichloroethylene	0.5	ND	0.5 ND
SURROGATES	SAI	MPLE	LAB BLANK
	AR	% REC.	AR % REC.
1,2-Dichloroethane-d4	40-140	122	40-140 125
Toluene-d8	64-139	104	64-139 102
BFB	50-149	109	50-149 108

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.)

ug/L = Micrograms Per Liter (ppb)

ND = Not Detected at or above the DLR. AR = Acceptable Range

• = DLR adjusted because of dilutions, concentrations, or limited sample.

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

Kelly A. Dunnahoo, B.S. Organic Laboratory Manager

KAD:cea

ANALYTICAL CHEMISTS

April 3, 1997

LAB No: SP 702023-6

Bermite Division of Whittaker 22116 W. Soledad Canyon Road

22116 W. Soledad Canyon Road Saugus , CA 91350 RE: Inorganic Analysis

Sample Site: MW5 Quarterly Sampling Area 317

Description: MW5/R/34

Sampled : March 19, 1997 Received : March 19, 1997 Completed : April 1, 1997

Sampled by : Abdun-Nur/Bricker Type of Sample: Monitoring Well

QA/QC ID# : 70202306-

Analytical Results

CONSTITUENT	METHOD	UNITS	DLR	RESULTS	MCL
Iron Manganese Sodium	200.7 200.8 200.7	ug/L ug/L mg/L	50 2.5* 1	60 ND 56	300 50

DLR = Detection Limit for Reporting Purposes. ND = Not Detected at or above the DLR.

ug/L = Micrograms Per Liter (ppb) mg/L = Milligrams Per Liter (ppm) mg/kg = Milligrams Per Kilogram

 ϕ = DLR adjusted because of dilutions, concentrations, or limited sample. Preservatives: (1) Cool 4°C (2) HN03 pH < 2 Containers: (a) Plastic

If you have any questions, please call.

FGL ENVIRONMENTAL

Kurt Wilkinson, B.S.

QA Director



April 3, 1997

INORGANIC Quality Assurance Report for sample: 702023

Bermite Division of Whittaker 22116 W. Soledad Canyon Road Saugus , CA 91350

	BATCH			BL	ANK QA/QC			CALIE	RATION	QA/QC					METHOD	QA/QC			
Constituent	ID	Method	Units	DLR	Result	NOTE	Туре	Conc.	% REC	AR	NOTE	Туре	Conc.	% REC	% REC	AR	% DIF	MAV	NOTE
Iron	2A 2A	200.7	ug/L	50	ND		ccv	5000	100	90-110		MS	500	99.0	99.3	75-125	0.3	20.0	
Manganese	13E 2A	200.8	ug/L-mg/L	0.50	ND		CCV	100	97.6	90-110		MS	80.0	90.8	91.2	75-125	0.3	20.0	
Sodium	2A 2A	200.7	mg/L	1.0	ND		CCV	25.0	97.6	90-110		MS	2.50	70.0	70.8	75-125	0.0	20.0	405
Chloride	8A 2C	4110B	mg/L	1.0	ND		CCV	50.0	91.9	90-110		MS	400	108	99.7	85-111	5.7	4.1	410
Conductivity	2A 2A	2510B	umhos/c2	1.0	ND		CCV	1000	98.0	75-125		Dup	1160	N/A	N/A	N/A	0.4	0.8	
pH	OA 2A	4500-н в	units		N/A		CCV	8.00	99.8	90-110		Dup	7.75	N/A	N/A	N/A	0.5	1.4	
Sulfate	8A 2C	4110B	mg/L	1.0	ND		CCV	100	92.6	90-110		MS	400	96.8	90.9	80-120	6.1	20.0	

FGL ID = 19970319 ND => Not Detected at ar above DLR. DLR => Detection Limit for Reporting purposes. N/A => Not Applicable NOTE => See note indicated below.

Notes:

405 Matrix Spike (MS) not within the Acceptance Range (AR) and/or Relative Percent Difference (RPD) not within the Maximum Allowable Value (MAV) because of low concentrations of the spike or sample. Batch qualified based on the LCS, CCV or ICV recovery.

410 Matrix Spike (MS) not within Acceptance Range (AR) and/or Relative Percent Difference (RPD) not within Maximum Allowable Value (MAV). Batch qualified based on the LCS, CCV, or ICV recovery.

FGL ENVIRONMENTAL, INC.

KW:cea

Kurt Wilkinson, B.S., QA Director



April 3, 1997

ORGANIC Quality Assurance Report for sample: 702023

Bermite Division of Whittaker 22116 W. Soledad Canyon Road Saugus , CA 91350

	BATCH	EPA			CALIE	RATION	QA/QC					METHOD	QA/QC			
Constituent	ID	Method	Units	Туре	Conc.	% REC	AR	NOTE	Туре	Conc.	% REC	% REC	AR	% DIF	MAV	NOTE
1,2-Dichloroethane-d4	1H12A	624	ug/L	LCS	10.0	116	40-140		BS	10.0	119	121	40-140	1.0	30.0	
Toluene-d8	1H12A	624	ug/L	LCS	10.0	105	64-139		BS.	10.0	104	104	64-139	0.0	30.0	
BFB	1H12A	624	ug/L	LCS	10.0	111	50-149		BS	10.0	110	110	50-149	0.5	30.0	
Trichloroethylene	1H12A	624	ug/L	LCS	10.0	106	37-151		BS	10.0	108	108	37-151	0.2	23.0	
TOC	0A 2A	415.1	mg/L	LCS	55.0	89.1	75-125		MS	50.0	92.0	98.0	75-125	6.3	20.0	
тох	5A 2A	TOX	ug/L	CCV	10.0	91.0	74-116		MS	100	91.0	101	75-125	10.4	20.0	

FGL ID = 19970320 N/A => Not Applicable NOTE => See note indicated below.

KAD:cea

Kurt Wilkinson, B.S., QA Director

FGL ENVIRONMENTAL, INC.

ANALYTICAL CHEMISTS

April 3, 1997

LAB No: SP 702025-1

Bermite Division of Whittaker 22116 W. Soledad Canyon Road

RE: Inorganic Analysis

Saugus, CA 91350

Sample Site: MW6 Quarterly Sampling Area 317

Description: MW6/A/34

Sampled: March 19, 1997 Received: March 19, 1997

Sampled by : Abdun-Nur/Bricker Type of Sample: Monitoring Well

Completed: March 21, 1997

QA/QC ID# : 70202501-

Analytical Results

CONSTITUENT	METHOD	UNITS	DLR	RESULTS
Conductivity	2510B	umhos/cm2	1	550
pH	4500-H B	units		7.7

DLR = Detection Limit for Reporting Purposes. ND = Not Detected at or above the DLR.

ug/L = Micrograms Per Liter (ppb) mg/L = Milligrams Per Liter (ppm) mg/kg = Milligrams Per Kilogram

• = DLR adjusted because of dilutions, concentrations, or limited sample.

Preservatives: (1) Cool 4°C Containers: (a) Plastic

If you have any questions, please call.

FGL ENVIRONMENTAL

Kurt Wilkinson, B.S.

QA Director

ANALYTICAL CHEMISTS

April 3, 1997

LAB No: SP 702025-3

Bermite Division of Whittaker 22116 W. Soledad Canyon Road RE: Organic Analysis Matrix: Monitoring Well

Saugus , CA 91350

Project/Site: MW6 Quarterly Sampling Area 317

Sample Description: MW6/C/34
Sampled by : Abdun-Nur/Bricker
Container : Ambon Class TEE Can

Sampled: March 19, 1997 Received: March 19, 1997

Container : Amber Glass TFE-Cap

Extracted : N/A

Preservatives:

Analyzed : March 25, 1997 QA/QC ID# : SP 97032500A A

TOTAL ORGANIC HALOGENS

CONSTITUENT	EPA METHOD	UNITS	SAMPLE DLR	SAMPLE RESULTS	LAB BLANK DLR RESULTS
тох	9020	ug/L	5	ND	5 ND

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.)
ug/L = Micrograms Per Liter (ppb)

ND = Not Detected at or above the DLR.

= DLR adjusted because of dilutions, concentrations, or limited sample.

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

Kelly A. Dunnahoo, B.S. Organic Laboratory Manager

KAD:cea

Field Office Visalia, CA TEL: 209/734-9473

ANALYTICAL CHEMISTS

April 3, 1997

LAB No: SP 702025-4

Bermite Division of Whittaker 22116 W. Soledad Canyon Road

Saugus , CA 91350

RE: Inorganic Analysis

Sample Site: MW6 Quarterly Sampling Area 317

Description: MW6/H/34
Sampled by : Abdun-Nur/Bricker
Type of Sample: Monitoring Well

Sampled: March 19, 1997 Received: March 19, 1997 Completed: March 27, 1997

QA/QC ID# : 70202504-

Analytical Results

CONSTITUENT	METHOD	UNITS	DLR	RESULTS
Chloride	4110B	mg/L	1	73
Sulfate	4110B	mg/L	1	29

DLR = Detection Limit for Reporting Purposes. ND = Not Detected at or above the DLR.

ug/L = Micrograms Per Liter (ppb) mg/L = Milligrams Per Liter (ppm) mg/kg = Milligrams Per Kilogram

• = DLR adjusted because of dilutions, concentrations, or limited sample.

Preservatives: (1) Cool 4°C Containers: (a) Plastic

If you have any questions, please call.

FGL ENVIRONMENTAL

Kurt Wilkinson, B.S.

QA Director

ANALYTICAL CHEMISTS

April 3, 1997

LAB No: SP 702025-5

Bermite Division of Whittaker 22116 W. Soledad Canyon Road

RE: Organic Analysis Matrix: Monitoring Well

Saugus , CA 91350

Project/Site: MW6 Quarterly Sampling Area 317

Sample Description: MW6/0/34 Sampled by : Abdun-Nur/Bricker

: March 19, 1997 Sampled Received: March 19, 1997

Container : VOA

Extracted: N/A

Preservatives:

Analyzed : March 21, 1997 QA/QC ID# : SP 97032101H A

EPA METHOD 624

CONSTITUENT	SAMPLE	SAMPLE	LAB	BLANK
	DLR	RESULTS	DLR	RESULTS
	ug/L	ug/L	ug/L	ug/L
Trichloroethylene	0.5	ND	0.5	ND
SURROGATES	SAM	PLE	LAB B	LANK
	AR	% REC.	AR %	REC.
1,2-Dichloroethane-d4	40-140	123	40-140	125
Toluene-d8	64-139	103	64-139	102
BFB	50-149	107	50-149	108

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.) ug/L = Micrograms Per Liter (ppb) ND = Not Detected at or above the DLR. AR = Acceptable Range • = DLR adjusted because of dilutions, concentrations, or limited sample.

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

Kelly A. Dunnahoo, B.S. Organic Laboratory Manager

KAD:cea

ANALYTICAL CHEMISTS

April 3, 1997

LAB No: SP 702025-6

Bermite Division of Whittaker 22116 W. Soledad Canvon Road

RE: Inorganic Analysis

Saugus , CA 91350

Sample Site: MW6 Quarterly Sampling Area 317

Description: MW6/R/34

: March 19, 1997 Sampled

Sampled by : Abdun-Nur/Bricker Type of Sample: Monitoring Well Received: March 19, 1997 Completed: March 31, 1997

QA/QC ID# : 70202506-

Analytical Results

CONSTITUENT	METHOD	UNITS	DLR	RESULTS	MCL
Iron Manganese Sodium	200.7 200.8 200.7	ug/L ug/L mg/L	50 0.5 1	170 1.6 56	300 50

DLR = Detection Limit for Reporting Purposes. ND = Not Detected at or above the DLR.

ug/L = Micrograms Per Liter (ppb) mg/L = Milligrams Per Liter (ppm) mg/kg = Milligrams Per Kilogram

♦ = DLR adjusted because of dilutions, concentrations, or limited sample. Preservatives: (1) Cool 4°C (2) HNO3 pH < 2 Containers: (a) Plastic</p>

If you have any questions, please call.

FGL ENVIRONMENTAL

Kurt Wilkinson, B.S.

QA Director



April 3, 1997

INORGANIC Quality Assurance Report for sample: 702025

Bermite Division of Whittaker 22116 W. Soledad Canyon Road Saugus , CA 91350

	BATCH			BL	ANK QA/QC			CALIE	RATION	QA/QC					METHOD	QA/QC			
Constituent	ID	Method	Units	DLR	Result	NOTE	Туре	Conc.	% REC	AR	NOTE	Туре	Conc.	% REC	% REC	AR	% DIF	MAV	NOTE
Iron	2A 2A	200.7	ug/L	50	ND		CCA	5000	100	90-110		MS	500	99.0	99.3	75-125	0.3	20.0	
Manganese	12E 2A	200.8	ug/L-mg/L	0.50	ND		ccv	100	98.4	90-110		MS	80.0	94.0	90.2	75-125	3.9	20.0	
Sodium	2A 2A	200.7	mg/L	1.0	ND		CCV	25.0	97.6	90-110		MS	2.50	70.0	70.8	75-125	0.0	20.0	405
Chloride ,	8A.2C	4110B	mg/L'	1.0 ·	ND '		-CCV	50.0	91.9	90-110	•	MS	400	108	99.7	85-111	5.7	4.1	410
Conductivity	2A 2A	2510B	umhos/c2	1.0	ND		CCV	1000	98.0	75-125		Dup	1160	N/A	N/A	N/A	0.4	0.8	
рH	0A 2A	4500-Н В	units		N/A		CCV	8.00	99.8	90-110		Dup	7.75	N/A	N/A	N/A	0.5	1.4	
Sulfate	8A 2C	4110B	mg/L	1.0	ND		CCA	100	92.6	90-110		MS	400	96.8	90.9	80-120	6.1	20.0	

FGL ID = 19970319 ND => Not Detected at ar above DLR. DLR => Detection Limit for Reporting purposes. N/A => Not Applicable NOTE => See note indicated below.

Notes:

405 Matrix Spike (MS) not within the Acceptance Range (AR) and/or Relative Percent Difference (RPD) not within the Maximum Allowable Value (MAV) because of low concentrations of the spike or sample. Batch qualified based on the LCS, CCV or ICV recovery.

410 Matrix Spike (MS) not within Acceptance Range (AR) and/or Relative Percent Difference (RPD) not within Maximum Allowable Value (MAV). Batch qualified based on the LCS, CCV, or ICV recovery.

FGL ENVIRONMENTAL, INC.

KW:cea

Kurt Wilkinson, B.S., QA Director



April 3, 1997

ORGANIC Quality Assurance Report for sample: 702025

Bermite Division of Whittaker 22116 W. Soledad Canyon Road Saugus , CA 91350

	BATCH	EPA			CALIE	RATION	QA/QC					METHOD	QA/QC			
Constituent	ID	Method	Units	Туре	Conc.	% REC	AR	NOTE	Туре	Conc.	% REC	% REC	AR	% DIF	MAV	NOTE
1,2-Dichloroethane-d4	1H12A	624	ug/L	LCS	10.0	116	40-140		BS	10.0	119	121	40-140	1.0	30.0	
Toluene-d8	1H12A	624	ug/L	LCS	10.0	105	64-139		BS	10.0	104	104	64-139	0.0	30.0	
BFB	1H12A	624	ug/L	LCS	10.0	111	50-149		BS	10.0	110	110	50-149	0.5	30.0	
Trichloroethylene	1H12A	624	ug/L	LCS	10:0	106 "	37-151		B\$	10.0	108	108	37-151	0.2	23.0	
TOC	OA ZA	415.1	mg/L	LCS	55.0	89.1	75-125		MS	50.0	92.0	98.0	75-125	6.3	20.0	
тох	5A 2A	TOX	ug/L	CCA	10.0	91.0	74-116		MS	100	91.0	101	75-125	10.4	20.0	

FGL ID = 19970320 N/A => Not Applicable NOTE => See note indicated below.

KAD:cea

FGL ENVIRONMENTAL, INC.

Kurt Wilkinson, B.S., QA Director

ANALYTICAL CHEMISTS

April 3, 1997

LAB No: SP 702027-1

Bermite Division of Whittaker 22116 W. Soledad Canyon Road RE: Inorganic Analysis

Saugus, CA 91350

Sample Site: MW10 Quarterly Sampling Area 317

Description: MW10/A/34

Sampled: March 19, 1997 Received: March 19, 1997

Sampled by : Abdun-Nur/Bricker Type of Sample: Monitoring Well

Completed: March 21, 1997

QA/QC ID# : 70202701-

Analytical Results

CONSTITUENT	METHOD	UNITS	DLR	RESULTS
Conductivity	2510B	umhos/cm2	1	600
pH	4500-H B	units		7.8

DLR = Detection Limit for Reporting Purposes. ND = Not Detected at or above the DLR.

ug/L = Micrograms Per Liter (ppb) mg/L = Milligrams Per Liter (ppm) mg/kg = Milligrams Per Kilogram

♦ = DLR adjusted because of dilutions, concentrations, or limited sample.

Preservatives: (1) Cool 4°C Containers: (a) Plastic

If you have any questions, please call.

FGL ENVIRONMENTAL

Kurt Wilkinson, B.S.

QA Director

ANALYTICAL CHEMISTS

April 3, 1997

LAB No: SP 702027-2

Bermite Division of Whittaker 22116 W. Soledad Canyon Road

RE: Organic Analysis Matrix: Monitoring Well

Saugus, CA 91350

Project/Site: MW10 Quarterly Sampling Area 317

Sample Description: MW10/B/34 Sampled by : Abdun-Nur/Bricker Container : Amber Glass TFE-Cap

Sampled : March 19, 1997 Received: March 19, 1997

Extracted: N/A

Preservatives: H2SO4 pH < 2

Analyzed: March 20, 1997 QA/QC ID# : SP 97032000A A

TOTAL ORGANIC CARBON

CONSTITUENT	EPA METHOD	UNITS	SAMPLE DLR	SAMPLE RESULTS	LAB DLR	BLANK RESULTS
TOC	415.1	mg/L	0.5	ND	0.5	ND

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.) ND = Not Detected at or above the DLR. mg/L = Milligrams Per Liter (ppm)

• = DLR adjusted because of dilutions, concentrations, or limited sample.

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

Kelly A. Dunnahoo, B.S. Organic Laboratory Manager

KAD:cea

ANALYTICAL CHEMISTS

April 3, 1997

LAB No: SP 702027-3

Bermite Division of Whittaker 22116 W. Soledad Canyon Road

RE: Organic Analysis Matrix: Monitoring Well

Saugus , CA 91350

Project/Site: MW10 Quarterly Sampling Area 317

Sample Description: MW10/C/34
Sampled by: Abdun-Nur/Bricker

Sampled : March 19, 1997 Received : March 19, 1997

Container : Amber Glass TFE-Cap

Extracted: N/A

Preservatives:

Analyzed : March 25, 1997 QA/QC ID# : SP 97032500A A

TOTAL ORGANIC HALOGENS

CONSTITUENT	EPA METHOD	UNITS	SAMPLE DLR	SAMPLE RESULTS	LAB BLANK DLR RESULTS
TOX	9020	ug/L	5	ND	5 ND

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.)
ug/L = Micrograms Per Liter (ppb)
ND = Not Detected at or above the DLR.

• = DLR adjusted because of dilutions, concentrations, or limited sample.

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

Kelly A. Dunnahoo, B.S. Organic Laboratory Manager

KAD: cea

ANALYTICAL CHEMISTS

April 3, 1997

LAB No: SP 702027-4

Bermite Division of Whittaker 22116 W. Soledad Canyon Road

RE: Inorganic Analysis

Saugus , CA 91350

Sample Site: MW10 Quarterly Sampling Area 317

Description: MW10/H/34 Sampled by : Abdun-Nur/Bricker Type of Sample: Monitoring Well

Sampled : March 19, 1997 Received: March 19, 1997

Completed: March 27, 1997

QA/QC ID# : 70202704-

Analytical Results

CONSTITUENT	METHOD	UNITS	DLR	RESULTS
Chloride	4110B	mg/L	1	68
Sulfate	4110B	mg/L		38

DLR = Detection Limit for Reporting Purposes. ND = Not Detected at or above the DLR. ug/L = Micrograms Per Liter (ppb) mg/L = Milligrams Per Liter (ppm) mg/kg = Milligrams Per Kilogram ♦ = DLR adjusted because of dilutions, concentrations, or limited sample. Preservatives: (1) Cool 4°C Containers: (a) Plastic

If you have any questions, please call.

FGL ENVIRONMENTAL

Kurt Wilkinson, B.S.

OA Director

ANALYTICAL CHEMISTS

April 3, 1997

LAB No: SP 702027-5

Bermite Division of Whittaker 22116 W. Soledad Canyon Road RE: Organic Analysis Matrix: Monitoring Well

Saugus . CA 91350

Project/Site: MW10 Quarterly Sampling Area 317

Sample Description: MW10/0/34 Sampled: March 19, 1997 Sampled by: Abdun-Nur/Bricker Received: March 19, 1997

Container: VOA Extracted: N/A

Preservatives:

Analyzed: March 21, 1997
QA/QC ID#: SP 97032101H A

EPA METHOD 624

CONSTITUENT	SAMPLE	SAMPLE	LAB BLANK
	DLR	RESULTS	DLR RESULTS
	ug/L	ug/L	ug/L ug/L
Trichloroethylene	0.5	ND	0.5 ND
SURROGATES	SAI	MPLE	LAB BLANK
	AR	% REC.	AR % REC.
1,2-Dichloroethane-d4	40-140	115	40-140 125
Toluene-d8	64-139	103	64-139 102
BFB	50-149	107	50-149 108

DLR = Detection Limit for Reporting Purposes. MCL = Maximum Contaminant Level (--- indicates none determined.)

ug/L = Micrograms Per Liter (ppb)

ND = Not Detected at or above the DLR. AR = Acceptable Range

• = DLR adjusted because of dilutions, concentrations, or limited sample.

See attached for Quality Assurance report. If you have any questions, please call.

FGL ENVIRONMENTAL

Kelly A. Dunnahoo, B.S. Organic Laboratory Manager

KAD:cea

Field Office Visalia, CA TEL: 209/734-9473 -

ANALYTICAL CHEMISTS

April 3, 1997

LAB No: SP 702027-6

Bermite Division of Whittaker 22116 W. Soledad Canyon Road

RE: Inorganic Analysis

Saugus, CA 91350

Sample Site: MW10 Quarterly Sampling Area 317

Description: MW10/R/34

Sampled : March 19, 1997

Sampled by : Abdun-Nur/Bricker Type of Sample: Monitoring Well Received: March 19, 1997 Completed: March 31, 1997

QA/QC ID# : 70202706-

Analytical Results

CONSTITUENT	METHOD	UNITS	DLR	RESULTS	MCL
Iron Manganese Sodium	200.7 200.8 200.7	ug/L ug/L mg/L	50 0.5 1	70 1.1 86	300 50

DLR = Detection Limit for Reporting Purposes. ND = Not Detected at or above the DLR.

ug/L = Micrograms Per Liter (ppb) mg/L = Milligrams Per Liter (ppm) mg/kg = Milligrams Per Kilogram

♦ = DLR adjusted because of dilutions, concentrations, or limited sample. Preservatives: (1) Cool 4°C (2) HNO3 pH < 2 Containers: (a) Plastic

If you have any questions, please call.

FGL ENVIRONMENTAL

Kurt Wilkinson, B.S.

QA Director



April 3, 1997

INORGANIC Quality Assurance Report for sample: 702027

Bermite Division of Whittaker 22116 W. Soledad Canyon Road Saugus , CA 91350

	BATCH			BI	BLANK QA/QC			CALIE	RATION	QA/QC					METHOD	QA/QC			
Constituent	ID	Method	Units	DLR	Result	NOTE	Туре	Conc.	% REC	AR	NOTE	Туре	Conc.	% REC	% REC	AR	% DIF	MAV	NOTE
Iron	2A 2A	200.7	ug/L	50	ND		ccv	5000	100	90-110		MS	500	99.0	99.3	75-125	0.3	20.0	
Manganese	12E 2A	200.8	ug/L-mg/L	0.50	ND		CCV	100	98.4	90-110		MS	80.0	94.0	90.2	75-125	3.9	20.0	
Sodium	2A 2A	200.7	mg/L	1.0	ND		CCV	25.0	97.6	90-110		MS	2.50	70.0	70.8	75-125	0.0	20.0	405
Chloride .	8A 2C	4110B	mg/L	1.0	ND.		CCV	50.0	91.9	90-110		MS	400	108	99.7	85-111	5.7	4.1	410
Conductivity	2A 2À	2510B	umhos/c2	1.0	ND		CCV	1000	98.0	75-125		Dup	1160	N/A	N/A	N/A	0.4	0.8	
рH	OA 2A	4500-Н В	units		N/A		CCV	8.00	99.8	90-110		Dup	7.75	N/A	N/A	N/A	0.5	1.4	
Sulfate	8A 2C	4110B	mg/L	1.0	ND	.,	CCV	100	92.6	90-110		MS	400	96.8	90.9	80-120	6.1	20.0	

FGL ID = 19970319 ND => Not Detected at ar above DLR. DLR => Detection Limit for Reporting purposes. N/A => Not Applicable NOTE => See note indicated below.

Notes:

405 Matrix Spike (MS) not within the Acceptance Range (AR) and/or Relative Percent Difference (RPD) not within the Maximum Allowable Value (MAV) because of low concentrations of the spike or sample. Batch qualified based on the LCS, CCV or ICV recovery.

410 Matrix Spike (MS) not within Acceptance Range (AR) and/or Relative Percent Difference (RPD) not within Maximum Allowable Value (MAV). Batch qualified based on the LCS, CCV, or ICV recovery.

FGL ENVIRONMENTAL, INC.

KW:cea

Kurt Wilkinson, B.S., QA Director



April 3, 1997

ORGANIC Quality Assurance Report for sample: 702027

Bermite Division of Whittaker 22116 W. Soledad Canyon Road Saugus , CA 91350

	ВАТСН	EPA			CALIBRATION QA/QC					METHOD QA/QC						
Constituent	ID	Method	Units	Туре	Conc.	% REC	AR	NOTE	Туре	Conc.	% REC	% REC	AR	% DIF	MAV	NOTE
1,2-Dichloroethane-d4	1H12A	624	ug/L	LCS	10.0	116	40-140		BS	10.0	119	121	40-140	1.0	30.0	
Toluene-d8	1H12A	624	ug/L	LCS	10.0	105	64-139		BS	10.0	104	104	64-139	0.0	30.0	
BFB	1H12A	624	ug/L	LCS	10.0	111	50-149		BS	10.0	110	110	50-149	0.5	30.0	
Trichloroethylene	1H12A	624	ug/L	LCS	10.0	106	37-151		BS	10.0	108	108	37-151	0.2	23.0	
TOC	0A 2A	415.1	mg/L	LCS	55.0	89.1	75-125		MS	50.0	92.0	98.0	75-125	6.3	20.0	
TOX	5A 2A	TOX	ug/L	ccv	10.0	91.0	74-116		MS	100	91.0	101	75-125	10.4	20.0	

FGL ID = 19970320 N/A => Not Applicable NOTE => See note indicated below.

FGL ENVIRONMENTAL, INC.

KAD:cea

Kurt Wilkinson, B.S., QA Director

APPENDIX H STATISTICAL ANALYSES

TABLE H-1
THIRTY-FOURTH QUARTER SAMPLING EVENT

				Well No.	
Parameter	Units	Tolerance Limit	MW-5	MW-6	MW-10
Specific Conductance	μmhos/cm ²	791	530	550	600
pН		7.07/7.99	7.8	7.7	7.8
TOC	mg/l	2.90	< 0.5	< 0.5	<0.5
TOX	μg/l	52.2	<5	<5	<5
Chloride	mg/l	196	54	73	68
Sulfate	mg/l	103	30	29	38
TCE	μg/l	0.5ª	< 0.5	< 0.5	<0.5
Iron	μg/l	184	60	170	70
Manganese	μg/l	21.0	<2.5	1.6	1.1
Sodium	mg/l	60.5	56	56	86

Note: All tolerance limits are upper limits except pH which has both upper and lower limits.

^{*}Tolerance limit set at detection limit.

TABLE H-2

CONCENTRATIONS OF GROUND WATER MONITORING PARAMETERS
IN SAMPLES FROM BACKGROUND MONITORING WELL MW-1

ъ.,			Conductance*	TOC*	TOX*	Cl Cl	SO ₄ 2	TCE	Fe	Mn	Na (()
Date 10/04/88 ^b	Quarter 1	pH* 7.5	(µmhos/cm²) 598	(mg/l) <3	(µgfl) <100	(mg/l)	(mg/l) 11	(μg/l) <5	(µg/l)	(µg/l)	(mg/l)
01/25/89	2	7.48	572	2.4°	<100		22	- 13			
04/17/89	3	7.2	312	<3	<100		11				
07/27/89	4	7.48	500	2.4°	<100		13				
10/31/89	5	7.6	524	<3	<100	83	10				
01/25/90	6	7.4	570	<3	<100	85	16				
04/17/90	7	7.55	504	<4	<20	88	11				
07/17/90	8	8.28	530	<4	<20	82	10				
10/18/90	9	7.4	544	<1	75°	98	23				
01/29/91	10	7.5	573	1.4	<5	96	8				
04/23/91	11	7.68	559	1.8	<5	100	10				
07/19/91	12	7.33	575	1.2	<5	97	11				
10/08/91 ^d	_		_	-							
03/13/92	14	7.5	639	0.4°	<5	131	13				
04/21/92	15	7.5	643	< 0.5	<5	130	9				
07/29/92	16	7.55	660	<0.5	6.9	133	11				
10/20/92	17	7.5	676	<0.5	<5	138	10				
01/27/93	18	7.68	707	<0.5	<5	137	6				
06/09/93	19	7.5	715	<0.5	<5	134	9		250	<30	52
07/14/93	20								220	<30	46
08/11/93	20	-							60	<30	54
09/22/93	20	7.5	720	<0.5	9	161	13		100	<30	52
12/08/93	21	7.4	726	<0.5	<5	151	10		50	<30	57
03/10/94	22	7.5	730	<0.5	<5	150	10	< 0.5	200	<30	48
06/22/94	23	7.5	740	<0.5	<5	150	15	<0.5	150	<30	54
09/14/94	24	7.4	750	< 0.5	8	160	9	< 0.5	60	2.5	57
12/14/94	25	7.5	770	<0.5	<5	150	10	< 0.5	80	4	51
03/29/95	26	7.5	770	<0.5	<5	160	12	< 0.5	60	1.6	49
06/27/95	27	7.4	760	<0.5	10	170	13	< 0.5	50	2.8	45
09/12/95	28	7.5	780	< 0.5	6	160	12	<0.5	90	3	53
12/08/95	29	6.9	780	<0.5	<5	180	12	< 0.5	< 50	2.7	50
03/20/96	30	7.4	770	<0.5	<5	180	13	< 0.5	<50	2.1	51
06/26/96	31	7.7	760	<0.5	<5	150	13	< 0.5	<50	2.1	53
09/19/96	32	7.6	750	< 0.5	7	160	12	< 0.5	80	1.8	52
12/05/96	33	7.5	750	<0.5	<5	160	10	<0.5	<50	2.6	55
03/19/97	34	7.8	770	<0.5	< 5	170	10	< 0.5	100	3.2	53

^{*}Each value reported before 06/09/93 is the average result from four replicate samples. Beginning 06/09/93, reported values are for a single sample as replicate sampling was stopped.

^bSamples from 01/27/88, 07/29/88, 08/15/88, and 10/04/88 reported TCE at $< 5 \mu g/l$.

The replicates included a portion with results below the detection limit. The average was calculated after assigning a value of one-half the detection limit for the samples below the detection limit.

^dNot sampled because water elevation dropped below elevation of sampling pump intake.

TABLE H-3

CONCENTRATIONS OF GROUND WATER MONITORING PARAMETERS
IN SAMPLES FROM BACKGROUND MONITORING WELL MW-3

Date	O	pH*	Conductance* (µmhos/cm³)	TOC* (mg/l)	TOX*	Cl (==#)	SO ₄ ² (mg/l)	TCE (µg/l)	Fe (µg/l)	Mn (μg/l)	Na (mg/l)
10/04/88 ^b	Quarter 1	7.48	699	<3	(μg/l) 361.25	(mg/l)	73	<5	(μχ/ι)	,μ <u>μ</u> (1)	(111641)
01/25/89	2	7.73	664	<3	<100		74				
04/17/89	3	7.73	- 004	<3	<100		9				
07/27/89	4	7.5	661	<3	<100		65				
10/31/89	5	7.53	617	<3	<100	35	68				
01/25/90	6	7.18	641	7.1°	<100	36	74				
04/17/90	7	7.33	590	<4	<20	46	60				
07/17/90	8	8.23	589	<4	<20	39	67				
10/18/90	9	7.63	642	0.7°	<100	34	15				
01/29/91	10	7.28	656	2.2	<5	54	80				
04/23/91	11	7.55	629	2.0	<5	34	77				
07/19/91	12	7.23	633	1.3	<5	45	85				
10/09/91	13	7.65	642	< 0.5	<5	37	34				
03/13/92	14	7.45	648	0.6	3.3°	33	85	***			
04/21/92	15	7.5	644	<0.5	<5	37	81				
07/29/92	16	7.55	643	0.34°	<5	33	74				
10/20/92	17	7.55	641	<0.5	<5	34	67				
01/27/93	18	7.6	640	< 0.5	<5	30	69				
06/09/93	19	7.6	627	< 0.5	<5	28	70		50	<30	48
07/14/93	20						_		<50	<30	44
08/11/93	20								<50	<30	50
09/22/93	20	7.4	630	<0.5	<5	37	87		<50	<30	50
12/08/93	21	7.4	627	< 0.5	<5	35	72		<50	<30	54
03/10/94	22	7.4	620	<0.5	<5	31	74	< 0.5	<50	<30	47
06/22/94	23	7.6	630	<0.5	8 ^d	29	71	< 0.5	< 50	<30	53
09/14/94	24	7.5	630	<0.5	<5	31	80	< 0.5	<50	0.7	52
12/14/94	25	7.5	630	<0.5	<5	28	69	< 0.5	<50	<1	48
03/29/95	26	7.7	620	< 0.5	7	28	71	< 0.5	<50	0.8	49
06/27/95	27	7.6	620	< 0.5	7	32	76	< 0.5	<50	0.6	53
09/12/95	28	7.6	620	< 0.5	<5	34	73	< 0.5	<50	<1	53
12/06/95	29	7.5	620	< 0.5	<5	29	77	< 0.5	<50	< 0.5	54
03/20/96	30	7.6	610	< 0.5	<5	35	91	< 0.5	<50	< 0.5	57
06/26/96	31	7.9	600	<0.5	<5	31	78	< 0.5	<50	3.3	61
09/19/96	32	8.0	610	< 0.5	<5	30	71	< 0.5	<50	< 0.5	56
12/05/96	33	7.6	600	<0.5	<5	26	62	< 0.5	<50	< 0.5	56
03/19/97	34	7.6	600	<0.5	<5	29	71	< 0.5	<50	< 0.5	60

^{*}Each value reported before 06/09/93 is the average result from four replicate samples. Beginning 06/09/93, reported values are for a single sample as replicate sampling was stopped.

^bSamples from 02/17/88, 05/27/88, 07/19/88, 08/15/88, and 10/04/88 reported TCE at <5 μ g/l.

The replicates included a portion with results below the detection limit. The average was calculated after assigning a value of one-half the detection limit for the samples below the detection limit.

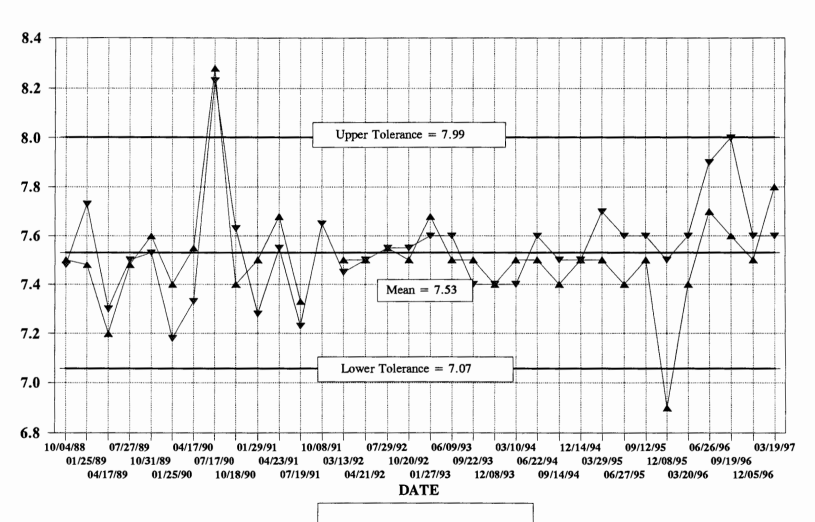
^dDuplicate sample analytical result also 8 μg/l.

TABLE H-4
TOLERANCE LIMIT CALCULATIONS

	pН	Conductance	TOC	TOX	Chloride	Sulfate	Iron	Manganese	Sodium
Σχ	504.5	42,188	55.84	884.70	4,964	2,738	2,125	246.05	1,878
n (number of samples)	67	65	67	66	59	67	36	36	36
x (mean)	7.53	649.05	0.83	13.40	84.14	40.87	59.03	6.83	52.2
s (sample standard deviation)	0.20	71.22	1.04	19.45	55.50	31.26	57.80	6.58	3.87
k (from tables)	2.309	2.000	1.994	1.997	2.021	1.994	2.158	2.158	2.158
Upper Tolerance Limit ^a	7.99	791	2.90	52.2	196	103	184	21.0	60.5
Lower Tolerance Limit ^b	7.07								

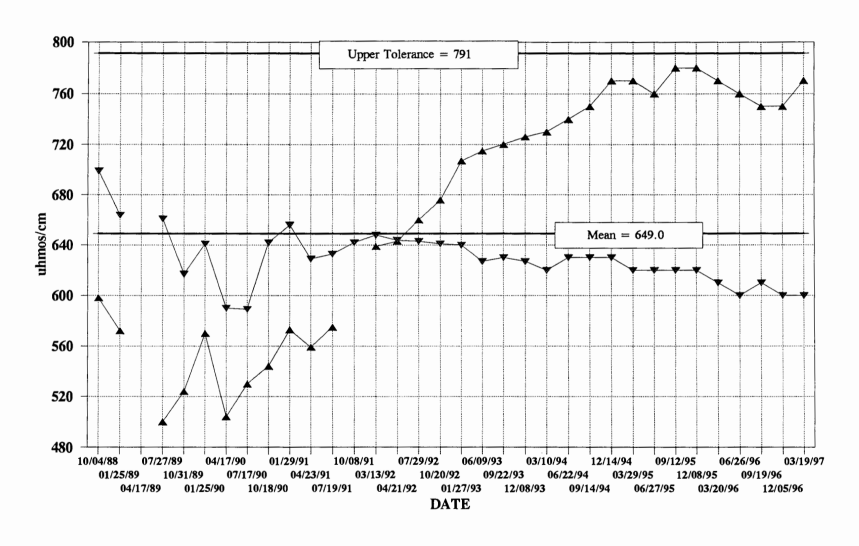
*Upper Tolerance Limit = \overline{x} + ks. *Lower Tolerance Limit = \overline{x} - ks.

pH MONITORING WELLS MW-1 AND MW-3

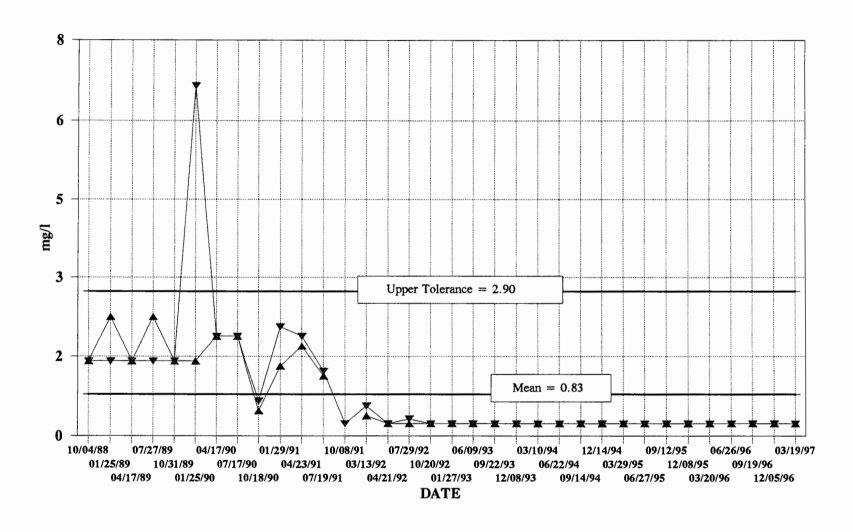


— MW-1 — MW-3

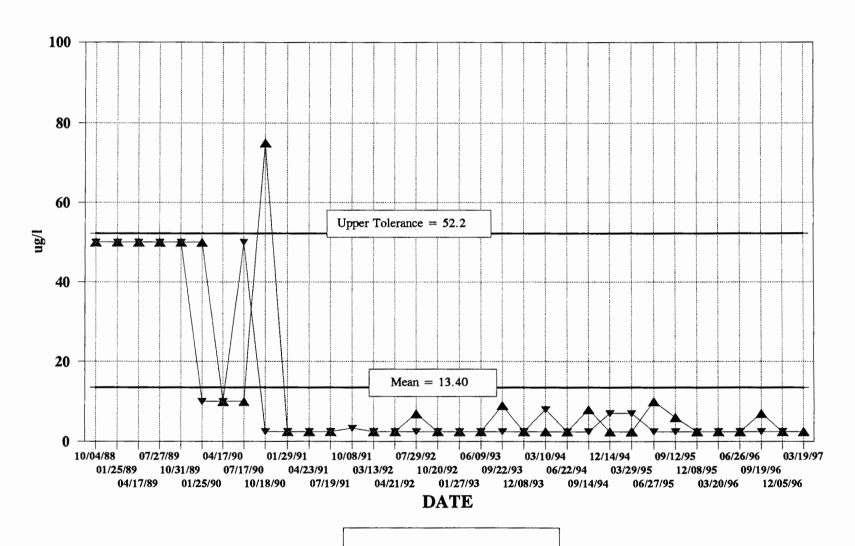
SPECIFIC CONDUCTANCE MONITORING WELLS MW-1 AND MW-3



TOTAL ORGANIC CARBON MONITORING WELLS MW-1 AND MW-3

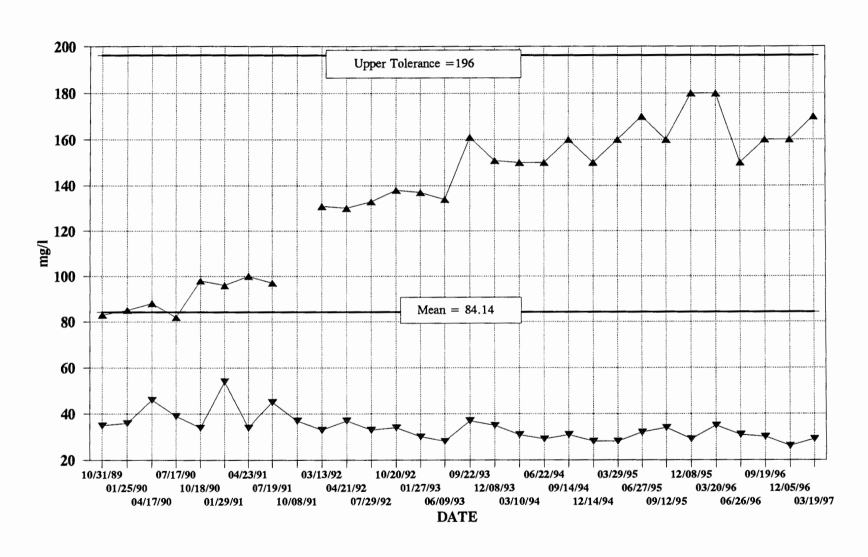


TOTAL ORGANIC HALOGENS MONITORING WELLS MW-1 AND MW-3



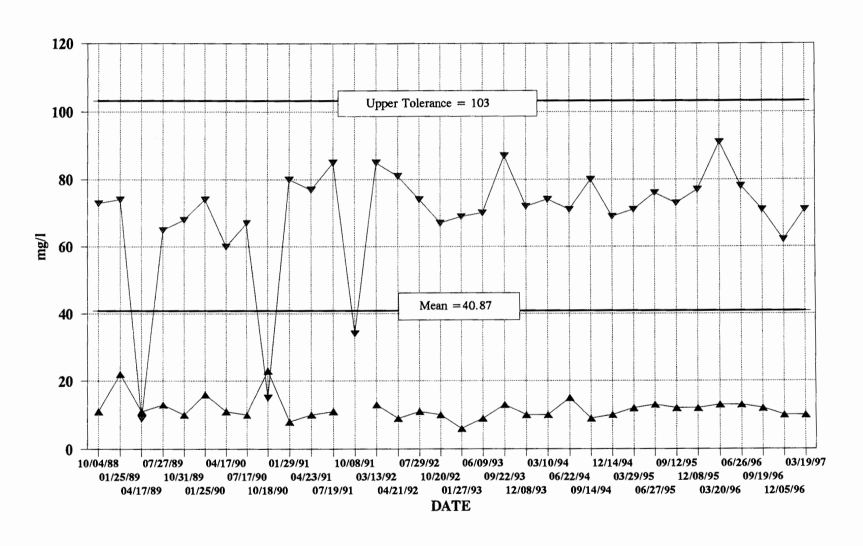
—<u>►</u> MW-1 — MW-3



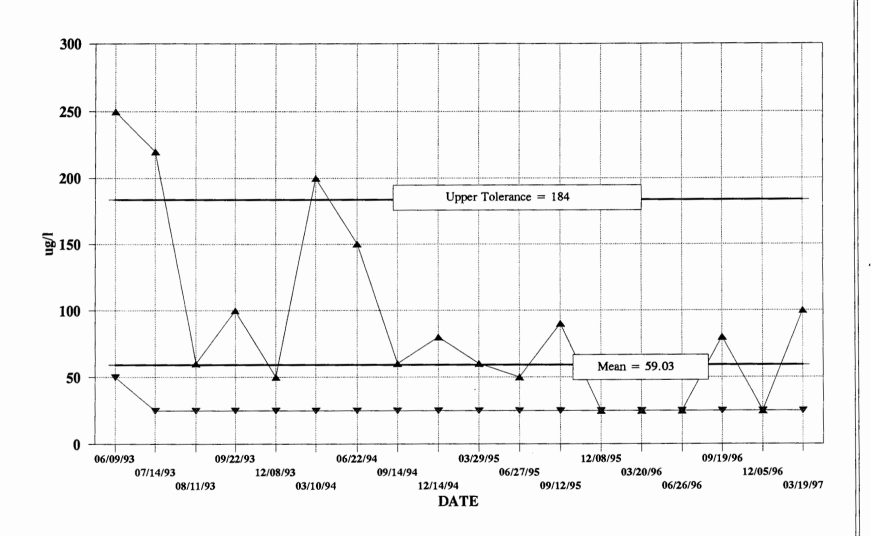


—<u>►</u> MW-1 —▼ MW-3

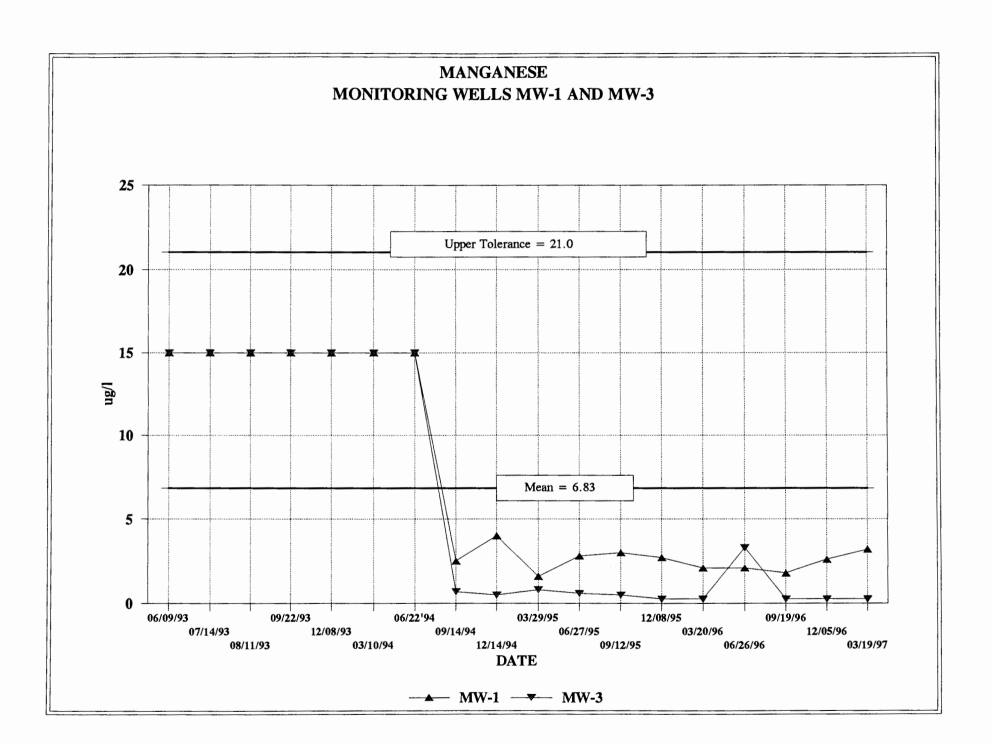


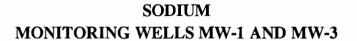


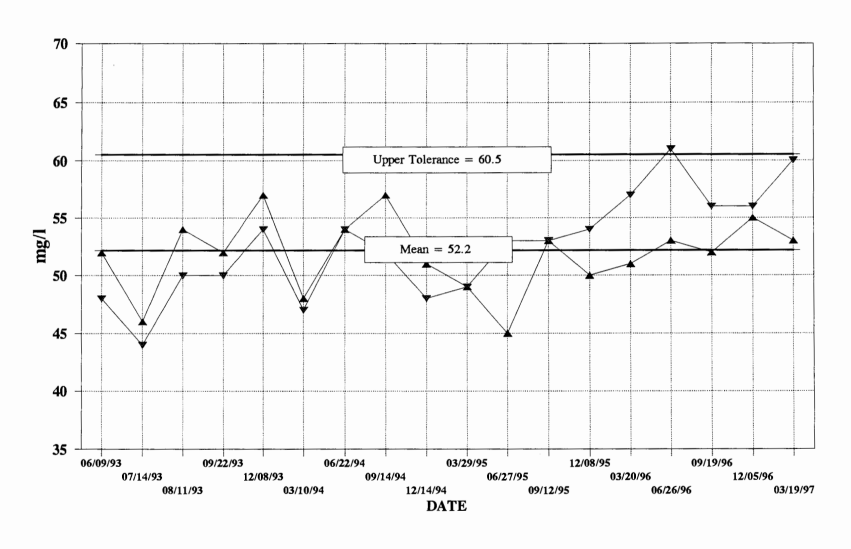




—<u>►</u> MW-1 —▼ MW-3







—— MW-1 —▼— MW-3